## Hector Reyes 10/623,237

#### => d his (FILE 'REGISTRY' ENTERED AT 12:06:23 ON 20 SEP 2004) DEL HIS Y E NATEGLINIDE/CN 1 S E3 L1STR 105816-04-4 L235 S L2 FUL FAM L3 SAVE L3 TEMP HECTOR/A FILE 'CAPLUS' ENTERED AT 12:07:34 ON 20 SEP 2004 302 S L3 L4110399 S POLYMORPH? L51165131 S CRYS? L6 11 S L4 AND L5 L724 S L4 AND L6 $^{\text{L8}}$ L9 24 S L7 OR L8 FILE 'BIOSIS, MEDLINE' ENTERED AT 12:10:15 ON 20 SEP 2004 351 S L3 L10233880 S CRYS? L114 S L10 AND L11 L12FILE 'USPATFULL' ENTERED AT 12:11:39 ON 20 SEP 2004 90 S L3 L13 12 S L13 AND (CRYS?)/TI,AB,CLM L14FILE 'CAPLUS, BIOSIS, MEDLINE, USPATFULL' ENTERED AT 12:12:23 ON 20 SEP 2004

35 DUP REM L9 L12 L14 (5 DUPLICATES REMOVED)

L15

=> fil caplus biosis medline uspatfull FILE 'CAPLUS' ENTERED AT 12:12:51 ON 20 SEP 2004 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

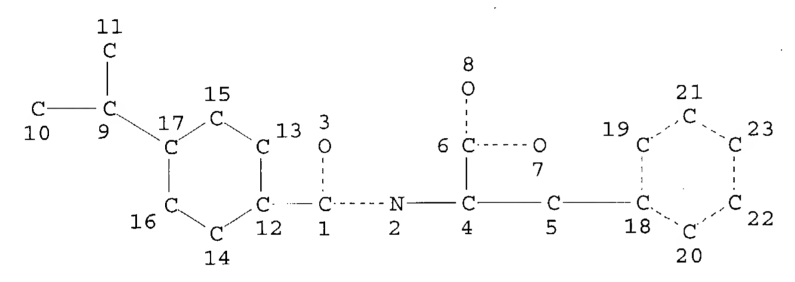
FILE 'BIOSIS' ENTERED AT 12:12:51 ON 20 SEP 2004 Copyright (c) 2004 The Thomson Corporation.

FILE 'MEDLINE' ENTERED AT 12:12:51 ON 20 SEP 2004

FILE 'USPATFULL' ENTERED AT 12:12:51 ON 20 SEP 2004
CA INDEXING COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)

=> d que 115

L2 STR



NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

DEFROIT DOBBVED 10 2111

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 23

STEREO ATTRIBUTES: NONE

35 SEA FILE=REGISTRY FAM FUL L2 L3302 SEA FILE=CAPLUS ABB=ON PLU=ON L3 L4110399 SEA FILE=CAPLUS ABB=ON PLU=ON POLYMORPH?/OBI L5PLU=ON CRYS?/OBI 1165131 SEA FILE=CAPLUS ABB=ON L611 SEA FILE=CAPLUS ABB=ON PLU=ON L4 AND L5 L724 SEA FILE=CAPLUS ABB=ON PLU=ON L4 AND L6 rs24 SEA FILE=CAPLUS ABB=ON PLU=ON L7 OR L8 Ь9 351 SEA L3 L10233880 SEA CRYS? L114 SEA L10 AND L11 L1290 SEA FILE=USPATFULL ABB=ON PLU=ON L3 L1312 SEA FILE=USPATFULL ABB=ON PLU=ON L13 AND (CRYS?)/TI,AB,CLM L1435 DUP REM L9 L12 L14 (5 DUPLICATES REMOVED) L15

=> d bib ab hitstr 115 1-35

L15 ANSWER 1 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2004:648496 CAPLUS

DN 141:179640

TI Preparation of a polymorphic crystalline form of the antidiabetic agent nateglinide

NPA

## Hector Reyes 10/623,237

```
Frenkel, Gustavo; Gome, Boaz; Wizel, Shlomit
IN
     Teva Pharmaceutical Industries Ltd., Israel; Teva Pharmaceuticals USA,
PA
     Inc.
SO
     PCT Int. Appl., 124 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN. CNT 3
     PATENT NO.
                                 DATE
                          KIND
                                             APPLICATION NO.
                                                                     DATE
                          A1
                                 20040812
                                             WO 2004-US839
                                                                     20040113
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     WO 2004067496
             AE, AE, AG, AL, AL, AM, AM, AM, AT, AT, AU, AZ, AZ, BA, BB, BG,
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             BG, BR, BR, BW, BY, BY, BZ, BZ, CA, CH, CN, CN, CO, CO, CR, CR,
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                                             WO 2003-US322375
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             GW, ML, MR, NE, SN, TD, TG
     US 2004181089
                          A1
                                 20040916
                                             US 2003-622905
                                                                     20030718
                           P
PRAI US 2003-442109P
                                 20030123
                           Ρ
                                 20030224
     US 2003-449791P
     US 2003-479016P
                           Ρ
                                 20030616
     US 2003-622905
                          A2
                                 20030718
     WO 2003-US22375
                          A2
                                 20030718
                          A2
                                 20031023
     US 2003-693166
                           A2
     US 2003-746697
                                 20031224
                                 20020718
     US 2002-396904P
                           Ρ
     US 2002-413622P
                           Ρ
                                 20020925
                                 20020926
     US 2002-414199P
     US 2002-423750P
                           P
                                 20021105
     US 2002-432093P
                           Р
                                 20021210
                           P
                                 20021212
     US 2002-432962P
                                 20030703
     US 2003-614266
                           Α
     The preparation of a polymorphic crystalline form (e.g., form U) of the
AB
antidiabetic
     agent nateglinide is described.
     105816-04-4, Nateglinide
IT
     RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES
     (Uses)
        (preparation of a polymorphic crystalline form of the
        antidiabetic agent nateglinide)
     105816-04-4 CAPLUS
RN
     D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI)
CN
     (CA INDEX NAME)
Absolute stereochemistry. Rotation (-).
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Ph R N CO<sub>2</sub>H O
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COPYRIGHT 2004 ACS on STN
     ANSWER 2 OF 35
                     CAPLUS
L15
     2004:203799 CAPLUS
AN
     140:241062
DN
     Process for the formation of a crystalline polymorphic
TI
     form of nateglinide
     Reguri, Buchi Reddy; Kadaboina, Rajasekhar; Polavarapu, Srinivas
IN
     Reddy's Laboratories Limited, India; Reddy's Laboratories, Inc.
PA
     PCT Int. Appl., 29 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
                          KIND
                                             APPLICATION NO.
                                                                     DATE
     PATENT NO.
                                             WO 2003-US26880
                                 20040311
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             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
             PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
             TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
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                                                                     20030827
     US 2004077725
                          A1
                                 20040422
                                             US 2003-649380
PRAI IN 2002-MA631
                                 20020828
                          Α
     A crystalline polymorphic form of nateglinide are described and its X-ray
AB
     diffraction pattern presented.
     105816-04-4P, Nateglinide
\operatorname{IT}
     RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
     (Reactant or reagent); USES (Uses)
        (process for the formation of a crystalline polymorphic
        form of nateglinide)
     105816-04-4 CAPLUS
RN
     D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI)
CN
     (CA INDEX NAME)
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# RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
ANSWER 3 OF 35 CAPLUS
L15
                              COPYRIGHT 2004 ACS on STN
AN
     2004:80637
                 CAPLUS
     140:151932
DN
     Preparation of polymorphic forms of nateglinide
TI
     Yahalomi, Ronit; Shapior, Evgeny; Dolitzky, Ben-zion; Gozlan, Yigael;
IN
     Gome, Boaz
     Teva Pharmaceutical Industries Ltd., Israel; Teva Pharmaceutical Usa, Inc
PA
SO
     PCT Int. Appl., 130 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 3
     PATENT NO.
                         KIND
                                 DATE
                                             APPLICATION NO.
                                                                     DATE
PΙ
     WO 2004009532
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                                 20040129
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     US 2004152782
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                                             US 2003-614266
                                                                     20030703
    US 2004116526
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     WO 2004067496
                          A1
                                 20040812
                                             WO 2004-US839
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             CU, CU, CZ, CZ, DE, DE, DK, DK, DM, DZ, EC, EC, EE, EE, EG, ES,
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             MZ, MZ, NA, NI
PRAI US 2002-396904P
                                 20020718
    US 2002-413622P
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                                 20020925
    US 2002-414199P
                          Ρ
                                 20020926
    US 2002-423750P
                          Ρ
                                 20021105
                          Ρ
    US 2002-432093P
                                 20021210
    US 2002-432962P
                          Ρ
                                 20021212
    US 2003-442109P
                          Ρ
                                 20030123
    US 2003-449791P
                          Ρ
                                 20030224
    US 2003~479016P
                          Ъ
                                 20030616
                          Α
    US 2003-614266
                                 20030703
    US 2002-393495P ·
                          P
                                 20020703
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US 2003-622905 A2 20030718 WO 2003-US22375 A2 20030718 US 2003-693166 A2 20031023 US 2003-746697 A2 20031224

The invention discloses the preparation of 26 characterized forms of nateglinide (forms A, C, D, F, G, I, J, K, L, M, N, O, P, Q, T, U, V, Y,  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$ ,  $\epsilon$ ,  $\sigma$ ,  $\theta$  and  $\Omega$ ). Most of the forms are solvates (with the exception of forms L,

P, U,  $\alpha$ ,  $\delta$  and  $\sigma$ ). Polymorphic forms are characterized by their mp, DSC, XRPD, FTIR; form interconversion is also discussed. For example, D-phenylalanine is reacted with trans-[[4-

(isopropyl)cyclohexanee]carbonyl]chloride (i. NaOHaq; ii. H2SO4). The wet cake of nateglinide is dissolved in EtOAc, the aqueous phase is removed and the resulting solution heated to  $50^{\circ}$  under reduced pressure and added to hot heptane. The resulting solution is cooled and seeded with the B-form to afford the  $\delta$ -form (33% yield).

IT 105816-04-4P, Nateglinide

RL: PEP (Physical, engineering or chemical process); PYP (Physical process); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); RACT (Reactant or reagent); USES (Uses)

(preparation of **polymorphic** forms of nateglinide)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

1T 105816-04-4DP, Nateglinide, polymorphs 651353-42-3P 651353-43-4P 651353-44-5P 651353-45-6P 651353-46-7P 651353-47-8P 651353-48-9P 651353-49-0P 651353-50-3P 651353-51-4P 651353-52-5P 651353-53-6P 651353-54-7P

RL: PEP (Physical, engineering or chemical process); PYP (Physical process); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses) (preparation of polymorphic forms of nateglinide)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

RN 651353-42-3 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with methanol (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

Absolute stereochemistry. Rotation (-).

CM 2

CRN 67-56-1 CMF C H4 O

HaC-OH

RN 651353-43-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with ethanol (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

CRN 64-17-5 CMF C2 H6 O

 $_{\mathrm{H_3C-CH_2-OH}}$ 

RN 651353-44-5 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with 1-butanol (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

Absolute stereochemistry. Rotation (-).

CM 2

CRN 71-36-3 CMF C4 H10 O

 $_{\rm H_3C^-CH_2^-CH_2^-OH}$ 

RN 651353-45-6 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with 1-propanol (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

CRN 71-23-8 CMF C3 H8 O

 $_{\mathrm{H_3C}-\mathrm{CH_2}-\mathrm{CH_2}-\mathrm{OH}}$ 

RN 651353-46-7 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with N,N-dimethylacetamide (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

Absolute stereochemistry. Rotation (-).

CM 2

CRN 127-19-5 CMF C4 H9 N O

Me | Me-N-Ac

RN 651353-47-8 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with 1-methyl-2-pyrrolidinone (9CI) (CA INDEX NAME)

CM ·1

CRN 105816-04-4 CMF C19 H27 N O3

CRN 872-50-4 CMF C5 H9 N O

RN 651353-48-9 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with N,N-dimethylformamide (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

Absolute stereochemistry. Rotation (-).

CM 2

CRN 68-12-2 CMF C3 H7 N O

RN 651353-49-0 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with 1,2-dimethoxyethane (9CI) (CA INDEX NAME)

CRN 105816-04-4 CMF C19 H27 N O3

Absolute stereochemistry. Rotation (-).

CM 2

CRN 110-71-4 CMF C4 H10 O2

 ${
m MeO-CH_2-CH_2-OMe}$ 

RN 651353-50-3 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with dimethylbenzene (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

Absolute stereochemistry. Rotation (-).

CM 2

CRN 1330-20-7 CMF C8 H10 CCI IDS



2 (D1-Me)

RN 651353-51-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with tetrachloromethane (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

Absolute stereochemistry. Rotation (-).

CM 2

CRN 56-23-5 CMF C Cl4

RN 651353-52-5 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with 1,2-dichloroethane (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

CRN 107-06-2 CMF C2 H4 Cl2

 ${\tt Cl-CH_2-CH_2-Cl}$ 

RN 651353-53-6 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with trichloromethane (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

Absolute stereochemistry. Rotation (-).

CM 2

CRN 67-66-3 CMF C H Cl3

RN 651353-54-7 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, compd. with heptane (9CI) (CA INDEX NAME)

CM 1

CRN 105816-04-4 CMF C19 H27 N O3

searched by Alex Waclawiw Page 13

Absolute stereochemistry. Rotation (-).

CM2

CRN 142-82-5 CMF C7 H16

 $Me^{-(CH_2)_5-Me}$ 

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 9 ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 35 USPATFULL on STN L152004:197476 USPATFULL ANProcess for preparing nateglinide and intermediates thereof TIYahalomi, Ronit, Kiryat Bialik, ISRAEL IN Shapiro, Evgeny, Haifa, ISRAEL Dolitzky, Ben-Zion, Petach Tiqva, ISRAEL Gozlan, Yigael, Ramot Sapir, ISRAEL US 2004152782 20040805 A1 PI20030703 (10) Α1 AIUS 2003-614266 20020703 (60) US 2002-393495P PRAI 20020718 (60) US 2002-396904P 20020925 (60) US 2002-413622P 20020926 (60) US 2002-414199P 20021105 (60) US 2002-423750P 20021210 (60) US 2002-432093P

Utility -DT

APPLICATION FS

KENYON & KENYON, ONE BROADWAY, NEW YORK, NY, 10004 LREP

20021212 (60)

20030123 (60)

20030224 (60)

Number of Claims: 57 CLMNExemplary Claim: 1 ECL 3 Drawing Page(s) DRWN

US 2002-432962P

US 2003-442109P

US 2003-449791P

LN.CNT 906

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

provided is a process for preparation of an intermediate in the ABsynthesis of nateglinide. Trans-4-isopropylcyclohexane acid chloride is formed by reacting 4-isopropylcyclohexanecarboxyl acid with thionyl chloride in the presence of an effective amount of an organic amide.

Also provided are processes for preparation of nateglinide by acylation of a suitable salt of D-phenylalanine with trans-4-isopropylcyclohexane acid chloride in both a single and a two phase system, and in water free of a co-solvent.

IT 105816-04-4P, Nateglinide

(process for preparation of nateglinide)

RN 105816-04-4 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

## IT 173653-89-9

(properties of nateglinide hydrate)

RN 173653-89-9 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, hydrate (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

### ●x H<sub>2</sub>O

ANSWER 5 OF 35 USPATFULL on STN L15 2004:185129 USPATFULL ANCombination of organic compounds TI Villhauer, Edwin Bernard, Morristown, NJ, UNITED STATES IN20040722 PΙ A1 US 2004143015 A1 20030910 (10) US 2003-471253 ΑI WO 2002-EP2665 20020311 Utility  $\mathtt{DT}$ FS APPLICATION NOVARTIS, CORPORATE INTELLECTUAL PROPERTY, ONE HEALTH PLAZA 430/2, EAST LREP HANOVER, NJ, 07936-1080 Number of Claims: 14 CLMNExemplary Claim: 1  $\mathsf{ECL}$ No Drawings DRWN LN.CNT 1075 CAS INDEXING IS AVAILABLE FOR THIS PATENT. The present invention relates to a combination of organic compounds ABwhich comprises at least two antidiabetic agents, preferably with different modes of action, in which the active ingredients are in each

case present in free form or in the form of a pharmaceutically

searched by Alex Waclawiw Page 15

acceptable salt and, optionally, at least on pharmaceutically acceptable carrier, for simultaneous, separate or sequential use.

IT 105816-04-4, Nateglinide

(pharmaceutical compns. containing combination of antidiabetic compds.)

RN 105816-04-4 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

```
ANSWER 6 OF 35 USPATFULL on STN
L15
       2004:152304 USPATFULL
AN
       Polymorphic forms of nateglinide
TI
       Yahalomi, Ronit, Kiryat Bialik, ISRAEL
IN
       Shapiro, Evgeny, Haifa, ISRAEL
       Dolitzky, Ben-Zion, Petach Tiqva, ISRAEL
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PRAI
       US 2002-396904P
       US 2002-413622P
                             20020925 (60)
                             20020926 (60)
       US 2002-414199P
                             20021105 (60)
       US 2002-423750P
                             20021210 (60)
       US 2002-432093P
                             20021212 (60)
       US 2002-432962P
                             20030123 (60)
       US 2003-442109P
                             20030224 (60)
       US 2003-449791P
                             20030616 (60)
       US 2003-479016P
       Utility.
\operatorname{DT}
       APPLICATION
FS
       KENYON & KENYON, ONE BROADWAY, NEW YORK, NY, 10004
LREP
       Number of Claims: 55
CLMN
       Exemplary Claim: 1
\mathsf{ECL}
       64 Drawing Page(s)
DRWN
LN.CNT 1830
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Provides are crystalline forms of nateglinide, labeled Forms
AB
       A, C, D, F, G, I, J, K, L, M, N, O, P, Q, T, U, V, Y, \alpha, \beta,
       \gamma, \delta, \epsilon, \sigma, 0 and \Omega, processes for
       their preparation and processes for preparation of other
       crystalline forms of nateglinide. Also provided are their
       pharmaceutical formulations and methods of administration.
    105816-04-4P, Nateglinide
IT
         (process for preparation of nateglinide)
     105816-04-4 USPATFULL
RN
     D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI)
CN
        (CA INDEX NAME)
```

IT 173653-89-9

(properties of nateglinide hydrate)

RN 173653-89-9 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, hydrate (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

●x H<sub>2</sub>O

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ANSWER 7 OF 35 USPATFULL on STN
L15
       2004:101855 USPATFULL
AN
       Crystalline form of N-(trans-4-isopropylcyclohexane
TI
       carbonyl) -D-phenylalanine and process for preparation thereof
       Reguri, Buchi Reddy, Hyderabad, INDIA
IN
       Kadaboina, Rajasekhar, Hyderabad, INDIA
       Polavarapu, Srinivas, Hyderabad, INDIA
       DR. REDDY'S LABORATORIES LIMITED (non-U.S. corporation)
PA
       DR. REDDY'S LABORATORIES, INC. (non-U.S. corporation)
                               20040422
       US 2004077725
                          A1
PI
                               20030827 (10)
ΑI
                          A1
       US 2003-649380
                           20020828
       IN 2002-6312002
PRAI
       Utility
DT
       APPLICATION
FS
       Janet I. Cord, Ladas & Parry, 26 West 61 Street, New York, NY, 10023
LREP
       Number of Claims: 33
CLMN
       Exemplary Claim: 1
ECL
       2 Drawing Page(s)
DRWN
LN.CNT 863
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       A new crystalline form of nateglinide is provided. The new
AB
       crystalline form is described by X-ray powder diffraction.
       Processes for making the new crystalline form of nateglinide
       are also provided.
    105816-04-4P, Nateglinide
IT
        (process for the formation of a crystalline polymorphic form of nateglinide)
     105816-04-4 USPATFULL
RN
```

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

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ANSWER 8 OF 35 USPATFULL on STN
L15
       2004:39640 USPATFULL
AN
       Methods for producing nateglinide crystals
TI
       Takahashi, Daisuke, Yokkaichi-Shi, JAPAN
IN
       Nishi, Seiichi, Yokkaichi-Shi, JAPAN
       Takahashi, Satoji, Yokkaichi-Shi, JAPAN
       AJINOMOTO CO. INC., Tokyo, JAPAN (non-U.S. corporation)
PA
                               20040212
                          A1
       US 2004030182
PI
                               20030418 (10)
       US 2003-418105
                          A1
AI
       Continuation of Ser. No. WO 2001-JP9069, filed on 16 Oct 2001, UNKNOWN
RLI
                           20001018
       JP 2000-317604
PRAI
       Utility
DT
       APPLICATION
FS
       OBLON, SPIVAK, MCCLELLAND, MAIER & NEUSTADT, P.C., 1940 DUKE STREET,
LREP
       ALEXANDRIA, VA, 22314
       Number of Claims: 13
CLMN
       Exemplary Claim: 1
\mathsf{ECL}
       No Drawings
DRWN
LN.CNT 387
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       There is provided methods for producing nateglinide crystals,
AB
       which comprises the steps of adding an acid(s) to a reaction mixture
       containing nateglinide to make it acidic, the reaction mixture being
       obtained by reacting trans-4-isopropylcyclohexylcarbonyl chloride with
       D-phenylalanine in a mixed solvent of ketone solvent and water in the
       presence of an alkali; and then adjusting the temperature of the mixture
       to 58° C. to 72° C. and the concentration of ketone
       solvent to more than 8 wt % and less than 22 wt % to conduct
       precipitation of nateglinide crystals. This producing method
       is the industrially beneficial methods for crystallization of
       nateglinide.
    105816-04-4P, Nateglinide
        (process for producing nateglinide crystals)
     105816-04-4 USPATFULL
RN
     D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI)
CN
       (CA INDEX NAME)
```

L15 ANSWER 9 OF 35 USPATFULL on STN

AN 2004:39426 USPATFULL

TI Nateglinide-containing hydrophilic pharmaceutical preparation

IN Ninomiya, Nobutaka, Kawasaki-Shi, JAPAN Makino, Chisato, Kawasaki-Shi, JAPAN Yabuki, Akira, Kawasaki-Shi, JAPAN

PA AJINOMOTO CO. INC, Tokyo, JAPAN (non-U.S. corporation)

PI US 2004029968 A1 20040212

AI US 2003-420886 A1 20030423 (10)

RLI Continuation of Ser. No. WO 2001-JP9292, filed on 23 Oct 2001, UNKNOWN

PRAI JP 2000-324374 20001024

DT Utility

FS APPLICATION

LREP OBLON, SPIVAK, MCCLELLAND, MAIER & NEUSTADT, P.C., 1940 DUKE STREET, ALEXANDRIA, VA, 22314

CLMN Number of Claims: 16

ECL Exemplary Claim: 1
DRWN 8 Drawing Page(s)

LN.CNT 486

AB

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

There is provided a nateglinide-containing hydrophilic pharmaceutical preparation comprising nateglinide B-type crystals as an effective ingredient, the contact angle of the surface of said preparation to water becoming 111 degree or less by incorporating in said preparation at least one hydrophilic substance selected from the groups consisting of hydrophilic polymers, surfactants, sugars, sugar alcohols and salts. This preparation is one having sufficient immediate-release and high dissolution properties, and can be easily prepared.

IT 105816-04-4, Nateglinide

(hypoglycemic hydrophilic drug prepns. containing)

RN 105816-04-4 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 10 OF 35 USPATFULL on STN

AN 2004:19511 USPATFULL

searched by Alex Waclawiw Page 19

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Hector Reyes 10/623,237
```

TI Nateglinide-containing preparation

IN Ninomiya, Nobutaka, Kawasaki-Shi, JAPAN

Makino, Chisato, Kawasaki-Shi, JAPAN

Yabuki, Akira, Kawasaki-Shi, JAPAN

PA AJINOMOTO CO. INC., Tokyo, JAPAN (non-U.S. corporation)

PI US 2004014815 A1 20040122

AI US 2003-421898 A1 20030424 (10)

RLI Continuation of Ser. No. WO 2001-JP9291, filed on 23 Oct 2001, UNKNOWN

PRAI JP 2000-324373 20001024

DT Utility

FS APPLICATION

LREP OBLON, SPIVAK, MCCLELLAND, MAIER & NEUSTADT, P.C., 1940 DUKE STREET, ALEXANDRIA, VA, 22314

CLMN Number of Claims: 20 ECL Exemplary Claim: 1

DRWN 9 Drawing Page(s)

LN.CNT 537

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention discloses, as a immediate-release preparation useful as an antidiabetic, a nateglinide-containing preparation comprising nateglinide as an active ingredient wherein the nateglinide is amorphous.

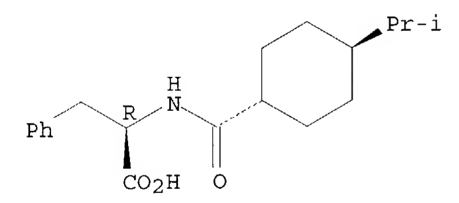
IT 105816-04-4, Nateglinide

(antidiabetic solid prepns. containing amorphous nateglinide and hydrophilic carriers)

RN 105816-04-4 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).



L15 ANSWER 11 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2003:892741 CAPLUS

DN 139:369757

Process for the preparation of a **crystal polymorphic** form of N-(trans-4-isopropylcyclohexylcarbonyl)-D-phenylalanine (nateglinide)

IN Rajamahendra, Shanmughasamy; Aswathanarayanappa, Chandrashekar; Puthiaparampil, Tom Thomas; Sridharan, Madhavan; Ganesh, Sambasivam

PA Biocon India Limited, India

SO PCT Int. Appl., 19 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI WO 2003093222 Al 20031113 WO 2002-IN114 20020429

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM,

#### Hector Reyes 10/623,237

HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG 20020429 PRAI WO 2002-IN114 Novel polymorph Form C of N-(trans-4-isopropylcyclohexylcarbonyl)-D-ABphenylalanine (I; i.e., nateglinide) is produced having a different IR spectrum and X-ray diffraction patterns (presented) from previously known forms of I. 105816-04-4P, Nateglinide ITRL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical process); PRP (Properties); PYP (Physical process); PREP (Preparation); PROC (Process) (process for the preparation of a crystal polymorphic form of N-(trans-4-isopropylcyclohexylcarbonyl)-D-phenylalanine (nateglinide)) 105816-04-4 CAPLUS RND-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI)

Absolute stereochemistry. Rotation (-).

(CA INDEX NAME)

## RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 12 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN 2003:837030 CAPLUS AN139:341723 DNNovel nateglinide crystals TIKoguchi, Yoshihito; Nakao, Tomoko; Sumikawa, Michito INPAAjinomoto Co., Inc., Japan SO PCT Int. Appl., 17 pp. CODEN: PIXXD2 DTPatent LAJapanese FAN.CNT 1 PATENT NO. KIND DATE

APPLICATION NO. DATE WO 2003-JP4686 **A**1 20031023 20030414 WO 2003087039 ΡĮ AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC,

## Hector Reyes 10/623,237

NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

PRAI JP 2002-111963 A 20020415

AB A type crystal (powder X-ray diffraction main peaks: 4.4°, 5.2°, 15.7°, 18.5° (2 theta)), M type crystal (powder X-ray diffraction main peaks: 6.0°, 14.2°, 15.2°, 18.8° (2 theta)), and P type crystal (powder X-ray diffraction main peaks: 4.8°, 5.3°, 14.3°, 15.2° (2 theta)) of nateglinide, which are all novel crystals, can be prepared by a method comprising dissolving nateglinide in a solvent exhibiting high solubility for nateglinide and then adding a solvent exhibiting poor solubility for nateglinide or dissolving nateglinide in a mixed solvent comprising a solvent exhibiting high solubility for nateglinide and a solvent exhibiting poor solubility for nateglinide and then cooling the resulting nateglinide solution to precipitate crystals, subjecting the product to filtration, and

drying at a specific temperature Nateglinide is a known antidiabetic.

IT 105816-04-4P, Nateglinide

RL: PRP (Properties); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of A, M, and P type nateglinide crystals by crystallization from mixture of solvents)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 13 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2003:837029 CAPLUS

DN 139:328379

TI 'Crystal polymorphism of nateglinide

IN Sutton, Paul Allen

PA Novartis A.-G., Switz.; Novartis Pharma G.m.b.H.

SO PCT Int. Appl., 10 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

	PATENT NO.						D	DATE		Ž	APPLICATION NO.						DATE			
				- <b></b> -		<b>~</b> ·	_						<del>-</del>							
ΡI	WO 2003087038					A1 20031023				7	WO 2	003-1		20030414						
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,		
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,		
			HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LT,	LU,		
			LV,	MA,	MD,	MK,	MN,	MX,	NI,	NO,	NZ,	OM,	PH,	PL,	PT,	RO,	RU,	SC,		
			SE,	SG,	SK,	TJ,	TM,	TN,	TR,	TT,	UA,	US,	UZ,	VC,	VN,	YU,	ZA,	ZW,		
			AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM									

RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR

PRAI US 2002-372625P P 20020415

AB New crystal forms of N-(trans-4-isopropylcyclohexylcarbonyl)-Dphenylalanine (i.e., nateglinide) are produced by dissolving nateglinide
in any of its forms, including solvates, in an organic solvent to form a
solution followed by precipitation of nateglinide from the solution, and
isolating and

drying the precipitated crystal form of nateglinide. The precipitation of nateglinide

may be induced either by cooling the solution, or by addition of another solvent

which is miscible with the first solvent but in which nateglinide is only poorly soluble, or by combination of the two. Depending on the solvent a specific crystal form of nateglinide may be obtained, e.g., the R'-type crystal form of nateglinide produced by the described method has a different m.p., infra red spectra and X-ray diffraction patterns from the previously known crystal forms of nateglinide.

IT 105816-04-4, Nateglinide

RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP (Physical process); PROC (Process)

(crystal polymorphism of nateglinide)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 14 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2003:221492 CAPLUS

DN 138:243310

TI Novel stable crystal form of N-trans-4-

isopropylcyclohexylcarbonyl) -D-phenylalanine and process of preparation

IN Shah, Vrajesh; Hitkari, Anurag; Deo, Keshav; Rengaraju, Srinivasan

PA Alembic Limited, India

SO PCT Int. Appl., 25 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1 KIND APPLICATION NO. DATE PATENT NO. DATE A120030320 WO 2001-IB2080 20011105 WO 2003022251 PΙ AE, AG, AL, AU, BA, BB, BG, BR, BZ, CA, CN, CO, CR, CU, CZ, DM, DZ, EC, EE, ES, GD, GE, HR, HU, ID, IL, IS, JP, KP, KR, LC, LK, LR, LT, LV, MA, MG, MK, MN, MX, NO, NZ, PH, PL, RO, SG, SI, SK, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,

DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG EP 2001-978760 20040714 **A**1 EP 1435912 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR PRAI IN 2001-MU871 20010912 Α A 20010912 IN 2001-MU872 20011105 W WO 2001-IB2080 A stable crystal form of N-(trans-4-isopropylcyclohexylcarbonyl)-D-ABphenylalanine (I) may be produced by crystallization of I with a solvent at 25

38 °C and forming crystals in the solvent. The crystal form may be formed by recrystn. out of solution The crystal form obtained in this way have different m.p., infra red spectrum and X-ray diffraction patterns

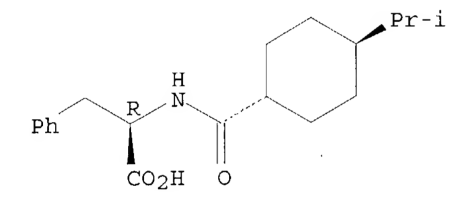
from previously known forms "B-type" and "H-Type" of the compound

(stable **crystal** form of N-trans-4-isopropylcyclohexylcarbonyl)-D-phenylalanine)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).



RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 15 OF 35 USPATFULL on STN L15 2003:325267 USPATFULL  $\mathbf{AN}$ Methods for producing nateglinide B-type crystals TI Sumikawa, Michito, Yokkaichi-Shi, JAPAN INMaruo, Makoto, Yokkaichi-Shi, JAPAN Miyazaki, Kazuo, Yokkaichi-Shi, JAPAN Nishina, Shigehiro, Yokkaichi-Shi, JAPAN Matsuzawa, Yukiko, Yokkaichi-Shi, JAPAN AJINOMOTO CO. INC, Tokyo, JAPAN (non-U.S. corporation) PA20031211 PIUS 2003229249 A1 20030424 (10) A1US 2003-421888 AIContinuation of Ser. No. WO 2001-JP9293, filed on 23 Oct 2001, UNKNOWN RLIJP 2000-324375 20001024 PRAI Utility DTFS APPLICATION OBLON, SPIVAK, MCCLELLAND, MAIER & NEUSTADT, P.C., 1940 DUKE STREET, LREP ALEXANDRIA, VA, 22314 Number of Claims: 9 CLMN Exemplary Claim: 1 ECL No Drawings DRWN LN.CNT 191

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A method for producing B-type crystals of nateglinide substantially free of H-type crystals is provided, which comprises drying solvated wet crystals of nateglinide at a low temperature until no solvent remains and making a crystal conversion thereof. According to this method, B-type crystals of nateglinide can be produced at an industrial scale without allowing other forms of the crystalline polymorphism to coexist.

IT 105816-04-4P, Nateglinide

(industrial process for producing B-form nateglinide crystals)

RN 105816-04-4 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

IT 173653-89-9

(industrial process for producing B-form nateglinide crystals)

RN 173653-89-9 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, hydrate (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

•x H<sub>2</sub>O

ANSWER 16 OF 35 USPATFULL on STN L15AN2003:232615 USPATFULL Method of treating metabolic disorders, especially diabetes, or a TIdisease or condition associated with diabetes Gatlin, Marjorie Regan, Hoboken, NJ, UNITED STATES INBall, Michele Ann, Morris Plains, NJ, UNITED STATES Mannion, Richard Owen, Mount Arlington, NJ, UNITED STATES Karnachi, Anees Abdulquadar, Hillsborough, NJ, UNITED STATES Guitard, Christiane, Hagenheim, FRANCE Allison, Malcolm, Basel, SWITZERLAND US 2003162816 20030828 PΙ Α1 US 2003-345908 20030116 (10) Α1 AI

RLI Continuation of Ser. No. US 2000-663264, filed on 15 Sep 2000, PENDING

PRAI GB 2000-21055 20000826

US 2000-304196P 20000407 (60)

US 2000-240918P 20000309 (60)

US 1999-240911P 19990917 (60)

DT Utility

FS APPLICATION

LREP THOMAS HOXIE, NOVARTIS, CORPORATE INTELLECTUAL PROPERTY, ONE HEALTH

PLAZA 430/2, EAST HANOVER, NJ, 07936-1080

CLMN Number of Claims: 41

ECL Exemplary Claim: 1
DRWN No Drawings

DRWN No Di LN.CNT 2226

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention relates to a combination, such as a combined preparation or pharmaceutical composition, respectively, which comprises nateglinide

(I) ##STR1##

or repaglinide and at least one other antidiabetic compound selected from the group consisting of thiazolidinedione derivatives (glitazones), sulfonyl urea derivatives and metformin for simultaneous, separate or sequential use in the prevention, delay of progression or treatment of diseases, especially metabolic disorders and in particular type 2 diabetes and diseases and conditions associated with diabetes; to a composition, respectively, which comprises nateglinide and a pharmaceutically acceptable carrier and to a process of making such composition; the use of such combination or composition for the preparation of a medicament for the prevention, delay of progression or treatment of diseases in warm-blooded animals; the use of such combination or composition for the cosmetic treatment of a mammal in order to effect a cosmetically beneficial loss of body weight; and to a method of improving the bodily appearance of a warm-blooded animal.

IT **105816-04-4**, Nateglinide

(pharmaceuticals containing nateglinide or repaglinide for treating diabetes or conditions associated with diabetes)

RN 105816-04-4 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 17 OF 35 USPATFULL on STN

AN 2003:93574 USPATFULL

TI Amino acid complexes of C-aryl glucosides for treatment of diabetes and method

IN Gougoutas, Jack Z., Princeton, NJ, UNITED STATES

PI US 2003064935 A1 20030403

US 6774112 B2 20040810

AI US 2002-117914 A1 20020408 (10)

PRAI US 2001-283097P 20010411 (60)

DT Utility

FS APPLICATION

LREP STEPHEN B. DAVIS, BRISTOL-MYERS SQUIBB COMPANY, PATENT DEPARTMENT, P O BOX 4000, PRINCETON, NJ, 08543-4000

CLMN Number of Claims: 19
ECL Exemplary Claim: 1

DRWN No Drawings

LN.CNT 1995

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Crystalline complexes are obtained from a 1:1 or 2:1 mixtures of either the (D) or (L) enantiomer of natural amino acids and compounds of formula ##STR1##

wherein

R.sup.1, R.sup.2 and R.sup.2a are independently hydrogen, OH, OR.sup.5, alkyl, --OCHF.sub.2, --OCF.sub.3, --SR.sup.5a or halogen;

R.sup.3 and R.sup.4 are independently hydrogen, OH, OR.sup.5b, alkyl, cycloalkyl, CF.sub.3, --OCHF.sub.2, --OCF.sub.3, halogen, --CONR.sup.6R.sup.6a, --CO.sub.2R.sup.5c, --CO.sub.2H, --COR.sup.6b, --CH(OH)R.sup.6c, --CH(OR.sup.5d)R.sup.6d, --CN, --NHCOR.sup.5e, --NHSO.sub.2R.sup.5f, --NHSO.sub.2Aryl, --SR.sup.5g, --SOR.sup.5h, --SO.sub.2R.sup.5i, or a five, six or seven membered heterocycle which may contain 1 to 4 heteroatoms in the ring which are N, O, S, SO, and/or SO.sub.2, or R.sup.3 and R.sup.4 together with the carbons to which they are attached form an annelated five, six or seven membered carbocycle or heterocycle which may contain 1 to 4 heteroatoms in the ring which are N, O, S, SO, and/or SO.sub.2;

R.sup.5, R.sup.5a, R.sup.5b, R.sup.5c, R.sup.5d, R.sup.5e, R.sup.5f, R.sup.5g, R.sup.5h and R.sup.5i are independently alkyl;

R.sup.6, R.sup.6a, R.sup.6b, R.sup.6c and R.sup.6d are independently hydrogen, alkyl, aryl, alkylaryl or cycloalkyl, or R.sup.6 and R.sup.6a together with the nitrogen to which they are attached form an annelated five, six or seven membered heterocycle which may contain 1 to 4 heteroatoms in the ring which are N, O, S, SO, and/or SO.sub.2.

A method is also provided for treating diabetes and related diseases employing an SGLT2 inhibiting amount of the above complex alone or in combination with another antidiabetic agent or other therapeutic agent. 105816-04-4, Nateglinide

(preparation of amino acid/C-aryl glucoside complexes for treatment of diabetes and related diseases)

RN 105816-04-4 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

L15 ANSWER 18 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2003:697592 CAPLUS

DN 140:187130

TI Study on stability of nateglinide polymorphism

AU Li, Gang; Xu, Qun Wei; Mo, Xiang Yin; Chen, Jia Ying; Su, Guo Qiang

CS Chemistry and Physics Centralaboratory, Nanjing Normal University, Nanjing, 210097, Peop. Rep. China

SO Chinese Chemical Letters (2003), 14(7), 730-733

CODEN: CCLEE7; ISSN: 1001-8417

PB Chinese Chemical Society

DT Journal

LA English

The stability of three forms of nateglinide, especially, S-form and H-form, was determined The S-form was a new crystal structure of nateglinide. Three forms of nateglinide were treated under different conditions such as in various temps., humidity, light, etc. Anal. of their crystal structures was performed by x-ray powder diffraction and their particle shapes were observed with scanning electron microscope. The results indicated that the stability of S-form of nateglinide is the best among the three forms and their particle shapes are quite different. The S-form is the sheet structure of layer upon layer, H-form looks like a hank of silk lines and the B-form is of clubbed shape.

IT 105816-04-4, Nateglinide

RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(stability of nateglinide polymorphs)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 19 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2003:146027 CAPLUS

DN 139:235199

TI Study on stability of nateglinide polymorphism

AU Li, Gang; Xu, Qun-Wei; Mo, Xiang-Yin; Chen, Jia-Ying; Su, Guo-Qiang

CS Testing & Analysis Center, Nanjing Normal University, Nanjing, 210097, Peop. Rep. China

SO Huaxue Xuebao (2003), 61(2), 291-294

CODEN: HHHPA4; ISSN: 0567-7351

PB Kexue Chubanshe

DT Journal

LA Chinese

AB A study has been made on the stability of three forms of nateglinide treated in different conditions, such as temperature, humidity, irradiation and so

searched by Alex Waclawiw Page 28

#### Hector Reyes 10/623,237

on. Anal. of the crystal structure was performed by x-ray powder diffraction. Their particle shapes were observed in scan electron microscope. The results show that the stability of S-form of nateglinide is the best among the three forms.

105816-04-4, Nateglinide IT

RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(stability of nateglinide polymorphism)

105816-04-4 CAPLUS RN

D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) CN(CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 20 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

2002:813874 CAPLUS AN

DN137:311199

Amino acid complexes of C-aryl glucosides for treatment of diabetes TI

Gougoutas, Jack Z. IN

Bristol-Myers Squibb Company, USA PA

PCT Int. Appl., 80 pp. SO

CODEN: PIXXD2

Patent DT

English  $_{
m LA}$ 

FAN.CNT 1																			
		PATENT NO.							DATE	DATE APPLICA					NO.		DATE		
														<del>-</del>					
P	Ί	WO	2002	0830	66		A2		20021024		1	WO 2	002-		20020408				
		WO	2002	0830	66		<b>A</b> 3		20030306										
			W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
				CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
	•			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,
										MG,									
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			RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	CH,
										GB,					-	-	•	•	•
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		US 2003064935								0403				-	·				
		US	6774	112			B2												•
		ΕP					A2		2004	0204	EP 2002-723801						20020408		
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Р	RAT	US	2001	•	·	•	•		•	•	,	,							
_			2002																
0	S						• •	•		0100									
V	U	MARPAT 137:311199																	

Crystalline complexes are obtained from 1:1 or 2:1 mixts. of either the (D) or AB (L) enantiomer of natural amino acids and compds. of formula I [R1, R2, R2a = H, OH, OR5, alkyl, OCHF2, OCF3, SR5a, halogen; R3, R4 = H, OH, OR5b,

alkyl, cycloalkyl, CF3, OCHF2, OCF3, halogen, CONR6R6a, CO2R5c, CO2H, COR6b, CH(OH)R6c, CH(OR5d)R6d, CN, NHCOR5e, NHSO2R5f, NHSO2-aryl, SR5q, SOR5h, SO2R5i, or a five, six or seven membered heterocycle which may contain 1 to 4 heteroatoms (N, O, S, SO, and/or SO2), or R3 and R4 together with the carbons to which they are attached form an annelated five, six or seven membered carbocycle or heterocycle which may contain 1 to 4 heteroatoms in the ring; R5, R5a-R5i are independently alkyl; R6, R6a-R6d are independently H, alkyl, aryl, alkylaryl or cycloalkyl, or NR6R6a form an annelated five, six or seven membered heterocycle which may contain 1 to 4 heteroatoms in the ring]. A method is also provided for treating diabetes and related diseases employing an SGLT2 (sodium dependent glucose transporters found in the intestine and kidney) inhibiting amount of the above complex alone or in combination with another antidiabetic agent or other therapeutic agent. Thus, I (R1 = 4-Me, R4 = 4-OCHF2, R2, R2a, R3 = H) was prepared by a multistep procedure starting from o-toluic acid, anisole, 2,3,4,6-tetra-O-benzyl-β-D-glucolactone, and CHF2Cl and treated with L-phenylalanine to form the crystalline 1:1 complex.

ma

IT 105816-04-4, Nateglinide

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of amino acid/C-aryl glucoside complexes for treatment of diabetes and related diseases)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 21 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2002:391524 CAPLUS

DN 136:374894

TI . Nateglinide-containing hydrophilic drug preparations

IN Ninomiya, Nobutaka; Makino, Chisato; Yabuki, Akira

PA Ajinomoto Co., Inc., Japan

SO PCT Int. Appl., 26 pp.

CODEN: PIXXD2

DT Patent

LA Japanese

FAN.CNT 1

	PATENT	KIN	D	DATE			APPL	ICAT		DATE								
			_						- ~									
PI	WO 2002040010				A1		2002	0523	1	WO 2001-JP9292						20011023		
	. W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,	
							DK,							-	•	•	-	
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	
							MD,											
		PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	
							ZW,											
	RW:						MZ,										CY,	
							GB,								•	•		

BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG 20020527 AU 2001-96000 AU 2001096000 A5 20011023 BR 2001014897 Α 20030812 BR 2001-14897 20011023 20030813 EP 1334721 **A**1 EP 2001-976818 20011023 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR A1 20040212 US 2003-420886 US 2004029968 20030423 PRAI JP 2000-324374 Α 20001024 WO 2001-JP9292 W 20011023

AB Hydrophilic drug prepns. contain nateglinide B crystals useful as a hypoglycemic agent as the active ingredient which comprises a hydrophilic substance selected from the group consisting of hydrophilic polymers, surfactants, sugars, sugar alcs. and salts, and thus have a contact angle of the preparation surface to water of 111° or less. These prepns., which are rapid release prepns. having high elution properties, can be easily produced.

IT 105816-04-4, Nateglinide

RL: BCP (Biochemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(hypoglycemic hydrophilic drug prepns. containing)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 22 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2002:332157 CAPLUS

DN 136:340998

TI Process for producing B-form nateglinide crystals

IN Sumikawa, Michito; Maruo, Makoto; Miyazaki, Kazuo; Nishina, Shigehiro; Matsuzawa, Yukiko

PA Ajinomoto Co., Inc., Japan

SO PCT Int. Appl., 9 pp.

CODEN: PIXXD2

DT Patent

LA Japanese

FAN.CNT 1

L WIM.																	
	PATENT	KIN	D	DATE		Ž	APPL	ICAT	DATE								
ΡI	WO 2002	A1		2002	0502	Ţ	WO 2	001-	20011023								
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	KZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PH,	PL,
		PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,
		US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	KΖ,	MD,	RU,	ΤJ,	MT	
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,

DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG AU 2001096001 **A5** 20020506 AU 2001-96001 20011023 EP 2001-976819 EP 1334964 **A**1 20030813 20011023 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR 20040225 BR 2001014846 Α BR 2001-14846 20011023 US 2003229249 **A**1 20031211 US 2003-421888 20030424 PRAI JP 2000-324375  $\mathbf{A}$ 20001024 WO 2001-JP9293 W 20011023

AB A process for producing B-form nateglinide crystals containing substantially no H-form crystals comprises the steps of drying wet crystals of a nateglinide solvate at a low temperature until the solvent disappears and then causing them to undergo a crystal transition. Nateglinide is a known antidiabetic. By this process, B-form nateglinide crystals can be produced on an industrial scale.

IT **105816-04-4P**, Nateglinide

RL: PAC (Pharmacological activity); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(industrial process for producing B-form nateglinide crystals

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

IT 173653-89-9

RL: PEP (Physical, engineering or chemical process); PROC (Process) (industrial process for producing B-form nateglinide crystals)

RN 173653-89-9 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, hydrate (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

●x H<sub>2</sub>O

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD

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ALL CITATIONS AVAILABLE IN THE RE FORMAT
L15
     ANSWER 23 OF 35 CAPLUS
                              COPYRIGHT 2004 ACS on STN
     2002:314896
AN
                  CAPLUS
DN
     136:325825
TI
     Process for producing nateglinide crystals
     Takahashi, Daisuke; Nishi, Seiichi; Takahashi, Satoji
IN
     Ajinomoto Co., Inc., Japan
PA
SO
     PCT Int. Appl., 14 pp.
     CODEN: PIXXD2
     Patent
\mathrm{DT}
LA
     Japanese
FAN.CNT 1
     PATENT NO.
                         KIND
                                 DATE
                                             APPLICATION NO.
PΙ
     WO 2002032854
                          Α1
                                 20020425
                                             WO 2001-JP9069
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL,
             PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG,
             US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     AU 2001094265
                          A5
                                 20020429
                                            AU 2001-94265
                                                                    20011016
                                20030813
                                            EP 2001-974875
     EP 1334963
                          A1
                                                                    20011016
           AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     BR 2001014729
                          Α
                                20031014
                                            BR 2001-14729
                                                                    20011016
     US 2004030182
                          A1
                                20040212
                                            US 2003-418105
                                                                    20030418
PRAI JP 2000-317604
                          A
                                20001018
     WO 2001-JP9069
                          W
                                20011016
     A process for producing nateglinide crystals comprises reacting
AB
     trans-4-isopropylcyclohexylcarbonyl chloride with D-phenylalanine in a
  · mixed solvent consisting of a ketone solvent and water in the presence of
     an alkali to obtain a reaction mixture containing nateglinide, adding an acid
to
    the reaction mixture to make it acidic, and regulating (a) the temperature to
     58° to 72° and (b) and the ketone solvent concentration to > 8 weight%
     and < 22 weight%, to conduct crystallization Nateglinide is a known
antidiabetic.
    The process is an industrially advantageous method for crystallizing
    nateglinide.
    105816-04-4P, Nateglinide
IT
    RL: IMF (Industrial manufacture); PRP (Properties); PUR (Purification or
    recovery); SPN (Synthetic preparation); THU (Therapeutic use); BIOL
     (Biological study); PREP (Preparation); USES (Uses)
        (process for producing nateglinide crystals)
     105816-04-4 CAPLUS
RN
    D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI)
CN
     (CA INDEX NAME)
```

Absolute stereochemistry. Rotation (-).

RE.CNT 3

## RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 24 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2003:686087 CAPLUS

DN 140:292376

TI Study on the crystal types of nateglinide

AU Sun, Piaoyang; Gou, Shaohua; Ma, Yonglin

CS State Key Laboratory of Coordination Chemistry, Nanjing University, Nanjing, 210093, Peop. Rep. China

SO 'Huaxue Yanjiu Yu Yingyong (2002), 14(4), 457-458, C3 CODEN: HYYIFM; ISSN: 1004-1656

PB Huaxue Yanjiu Yu Yingyong Bianjibu

DT Journal

LA Chinese

AB N-(trans-4-methylethylcyclohexylcarbonyl)-D-phenylalanine, nateglinide, is an effective drug to decrease blood sugar, which is under clin. trials in China. This compound has been reported to have two crystal types, one of which is more suitable to prepare the drug. The nateglide with different crystal types was prepared Their m.ps., TGA-DTA and DSC spectral data, LR and X-ray powder diffraction spectra of all samples were studied with different crystal types. A new crystal type that has not been reported in the literature was discovered. The method for controlling the crystal type was also presented.

IT 105816-04-4, Nateqlinide

RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(polymorphism; polymorphism of nateglinide)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 25 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2002:811385 CAPLUS

DN 139:12440

TI , Identification of nateglinide and its crystal forms in nateglinide tablets using IR Spectra subtraction techniques AU Lin, Kejiang; Chen, Wei; Tang, Weiguo; You, Qidong

searched by Alex Waclawiw Page 34

## Hector Reyes 10/623,237

- CS Department of Medicinal Chemistry, China Pharmaceutical University, Nanjing, 21009, Peop. Rep. China
- SO Zhongguo Yaoke Daxue Xuebao (2002), 33(2), 124-126 CODEN: ZHYXE9; ISSN: 1000-5048
- PB Zhongguo Yaoke Daxue
- DT Journal
- LA Chinese
- The innovational identification method of IR (eliminated method) for detection of the crystal form of nateglinide in prepns. was presented. The IR spectrum by spectra subtraction techniques was obtained by subtracting IR spectrum after adding small volume of solvent to eliminate nateglinide from the spectrum of nateglinide tablets' KBr disk to identify the crystal form of nateglinide. The method (eliminated method) was useful in identification of the nateglinide crystal form in prepns.
- nateglinide tablets using IR spectra subtraction techniques)
  RN 105816-04-4 CAPLUS
- CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

- L15 ANSWER 26 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2002:609152 CAPLUS
- DN 138:254901
- TI / a new synthesis method of nateglinide as antidiabetic drug
- AU Wang, Dun; Liang, Yiheng; Gong, Ping; Zhao, Yanfang
- CS School of Pharmaceutical Engineering, Shenyang Pharmaceutical University, Shenyang, 110016, Peop. Rep. China
- SO | Zhongguo Yaowu Huaxue Zazhi (2002), 12(2), 94-96
- CODEN: ZYHZEF; ISSN: 1005-0108
- PB Zhongguo Yaowu Huaxue Zazhi Bianjibu
- DT / Journal
- LA Chinese
- OS CASREACT 138:254901
- AB A new antidiabetic drug-nateglinide was synthesized from isopropylbenzene by Friedel-Crafts reaction, chloroform reaction, catalytic hydrogenation to obtain trans-4-isopropylhexanecarboxylic acid, acylation of D-phenylalanine Et ester, hydrolysis to obtain nateglinide B-type crystal, and crystal-conversion. The total yield was 9.8%.
- IT | 105816-04-4P, Nateglinide
  - RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
  - (synthesis of nateglinide as antidiabetic drug)
- RN 105816-04-4 CAPLUS
- CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 27 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 1

AN 2002:234892 CAPLUS

DN 137:39555

TI Detection of crystal polymorphs of nateglinide by DSC

AU Lin, Kejiang; Chen, Wei; You, Qidong

CS China Pharmaceutical University, Nanjing, 210009, Peop. Rep. China

SO Yaoxue Xuebao (2002), 37(1), 46-49 CODEN: YHHPAL; ISSN: 0513-4870

PB Yaoxue Xuebao Bianjibu

DT Journal

LA Chinese

AB

The differential scanning calorimetric (DSC) methodol. for controlling the crystal-type B form of nateglinide was presented. Pure fine powder of crystal-type B and H of nateglinide dried with P205 as desiccant at 80° in vacuum for 4 h was measured dQ/dT by DSC at heating rate of 10° min-1 and temperature between 100° and 200° to calculate the enthalpy  $\Delta HB$  and  $\Delta HH$ . Uniform mixts. of crystal-type B and H of dried fine powder of nateglinide in different proportions were accurately weighed. The enthalpy of the mixts. was measured by DSC as above to calculate the enthalpy ( $\Pi\Delta H$ ). Using B% as X,  $\Pi\Delta H$ as parameters, the regression equation was obtained. Based on this equation, the unknown composition of mixed crystal was evaluated by  $y\delta H$ values. The method was used to control the limitation of crystal-type B of nateglinide by the  $H\delta H$  value of mixture of known composition as reference The results measured from different labs. showed that the repeatability was 0.61% and recoveries were 86.2-127% when the amount of crystal-type B was between 0-15%. This method can be used to evaluate the crystal-type B composition of nateglinide.

IT 105816-04-4, Nateglinide

RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP (Physical process); PROC (Process)

(detection of crystal polymorphs of nateglinide by DSC)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

- L15 ANSWER 28 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2003:762699 CAPLUS
- DN 140:64875
- TI Study of nateglinide polymorphism
- AU Li, Gang; Xu, Qunwei; Yao, Jie; Su, Guoqiang; Wang, Fang
- CS Chemistry and Physics Central-laboratory, Nanjing Normal University, Nanjing, 210097, Peop. Rep. China
- SO Huagong Shikan (2002), 16(7), 17-18 CODEN: HUSHFT; ISSN: 1002-154X
- PB Huagong Shikan Zazhishe
- DT Journal
- LA Chinese
- The crystal structure of nateglinide called an S form determined by an x-ray powder diffraction method. The pattern, data, and crystal size were obtained. The m.p. was determined by DSC as 172.04°.
- IT 105816-04-4, Nateglinide
  RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
- (nateglinide polymorphism)
- RN 105816-04-4 CAPLUS
- CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

- L15 ANSWER 29 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN
- AN 2001:283772 CAPLUS
- DN 134:285620
- TI Method of treating metabolic disorders with nateglinide
- IN Gatlin, Marjorie Regan; Pongowski, Michele; Dunning, Beth
- PA Novartis A.-G., Switz.; Novartis-Erfindungen Verwaltungsgesellschaft m.b.H.
- SO PCT Int. Appl., 28 pp. CODEN: PIXXD2
- DT Patent
- LA English
- FAN.CNT 1

L MIA.	CIAT	Τ																
	PA	rent :	NO.			KIN	D	DATE			APPL	ICAT	ION :	NO.		D.	ATE	
				<b>-</b>			_					<b>-</b>				_		
PI	WO	2001	0266	39		A2		2001	0419		WO 2	000-	EP98	16		2	0001	006
	WO	2001	0266	39		<b>A</b> 3		2002	0110									
		W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,
			HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,
			LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
			SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,
			YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	KΖ,	MD,	RU,	TJ,	TM	·	•	•	•
		RW:						MZ,							AT.	BE.	CH.	CY.

DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

EP 1218015

A2 20020703

EP 2000-972695

20001006

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL

PRAI US 1999-415307

A 19991008

US 1999-415308

A 19991008

WO 2000-EP9816 W 20001006

The invention relates to a combination which comprises nateglinide and (a) an antidiabetic phenylacetic acid derivative or (b) acarbose for simultaneous, sep. or sequential use, in particular in the treatment of diseases, especially metabolic disorders; to a method of prevention, delay of progression or treatment of metabolic disorders, more especially diabetes, or a disease or condition associated with diabetes, and to a method of improving the bodily

IT 105816-04-4, Nateglinide
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (treating metabolic disorders with nateglinide)

RN 105816-04-4 CAPLUS
CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI)
(CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

appearance of a warm-blooded animal.

L15 ANSWER 30 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2002:130037 CAPLUS

DN 137:325603

TI Synthesis of Nateglinide

AU Zhu, Xue-yan; Peng, Ka; Wang, Xiao-qin; Yang, Li-ping

CS Dep. Chem., East China Normal Univ., Shanghai, 200062, Peop. Rep. China

SO Hecheng Huaxue (2001), 9(6), 537-540 CODEN: HEHUE2; ISSN: 1005-1511

PB Hecheng Huaxue Bianjibu

DT Journal

LA Chinese

OS \ CASREACT 137:325603

AB Title compound, a new antidiabetes medicine, was synthesized from iso-propylbenzene in seven steps, giving the product with overall yield 22%.

IT / 105816-04-4DP, Nateglinide, B crystal type

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and crystalline forms of)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

searched by Alex Waclawiw Page 38

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Mel

RL: SPN (Synthetic preparation); PREP (Preparation) (synthesis of Nateglinide

L15 ANSWER 31 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 2

AN 2001:625224 CAPLUS

DN 136:348527

TI New crystal form of nateglinide

AU Li, Gang; Su, Guoqiang; Xu, Qunwei; Zhu, Chongquan

CS Chemistry and Physics Central Laboratory, Nanjing Normal University, Nanjing, 210097, Peop. Rep. China

SO Yaoxue Xuebao (2001), 36(7), 532-534

CODEN: YHHPAL; ISSN: 0513-4870

PB Yaoxue Xuebao Bianjibu

DT Journal

LA | Chinese

The S form crystals of nateglinide [N-(trans-4-

isopropylcyclohexylcarbonyl)-D-phenylalanine] were studied by XRD, IR, elemental anal., and differential scan calorimetry. The S-form nateglinide crystal was different from the H-form or B-form. The m.p. was 172.04°. The results showed that the S-form nateglinide was a new

crystal form.

IT | 105816-04-4, Nateglinide

RL: PAC (Pharmacological activity); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(new crystal form of nateglinide)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 32 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 2001:844448 CAPLUS

DN 136:159110

TI A new **crystal** structure in nateglinide found by X-ray powder diffraction

AU Li, Gang; Su, Guo-qiang; Xu, Qun-wei

CS Center for Analysis & Measurement, Nanjing Normal University, Nanjing, 210097, Peop. Rep. China

SO Yaowu Fenxi Zazhi (2001), 21(5), 342-344

searched by Alex Waclawiw Page 39

CODEN: YFZADL; ISSN: 0254-1793 Yaowu Fenxi Zazhi Bianji Weiyuanhui PB Journal DTChinese LA A new crystal structure being assigned as S-form was found in nateglinide. AB The x-ray pattern and data were given and the m.p. was determined Phase anal. was carried out by x-ray powder diffraction; the m.ps. were determined by DSC. S-form nateglinide was different from the H or B crystal form. was 172.04°. S-form nateglinide was a new crystal form. X-ray powder diffraction anal. was one of the most effective methods for phase structure characterization. 105816-04-4, Nateglinide ITRL: PRP (Properties) (crystal structure of) 105816-04-4 CAPLUS RND-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) CN

(CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 33 OF 35 USPATFULL on STN AN96:9521 USPATFULL Crystals of N-(trans-4-isopropylcyclohexycarbonyl)-D-TIphenylalanine and methods for preparing them Sumikawa, Michito, Kawasaki, Japan INKoguchi, Yoshihito, Kawasaki, Japan Ohqane, Takao, Kawasaki, Japan Irie, Yasuo, Kawasaki, Japan Takahashi, Satoji, Yottukaichi, Japan Ajinomoto Co., Inc., Tokyo, Japan (non-U.S. corporation) PAPIUS 5488150 19960130 US 1993-166144  $\mathsf{AI}$ 19931214 (8) Continuation of Ser. No. US 1992-921224, filed on 29 Jul 1992, now RLIabandoned JP 1991-189696 PRAI 19910730 JP 1991-199453 19910808 Utility  $\operatorname{DT}$ FS Granted EXNAM Primary Examiner: Henley, III, Raymond; Assistant Examiner: MacMillan, Keith Oblon, Spivak, McClelland, Maier, & Neustadt LREP Number of Claims: 13 CLMN Exemplary Claim: 1 ECL 7 Drawing Figure(s); 5 Drawing Page(s) DRWN LN.CNT 528 CAS INDEXING IS AVAILABLE FOR THIS PATENT. Stable crystals of N-(trans-4-isopropylcyclohexylcarbonyl)-D-ABphenylalanine may be produced by treating this compound with a solvent at a temperature of at least 10° C. and forming crystals in the solvent at a temperature of at least 10° C. For example, searched by Alex Waclawiw Page 40

crystals may be formed by crystallization out of solution, or may be formed from solid particles of the compound suspended in a solvent. Crystals formed in this way have different melting point, infra red spectrum and X-ray diffraction patterns from previously known forms of the compound and have enhanced processability, eg. stability to grinding.

IT 105816-04-4P

(crystals, stable, preparation of)

RN 105816-04-4 USPATFULL

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L15 ANSWER 34 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN DUPLICATE 3

AN 1995:964992 CAPLUS

DN 124:155974

TI **Crystals** of N-(trans-4-isopropylcyclohexylcarbonyl)-D-phenylalanine and methods for preparing them

IN Sumikawa, Michito; Koguchi, Yoshihito; Ohgane, Takao; Irie, Yasuo; Takahashi, Satoji

PA Ajinomoto Co., Inc., Japan

SO U.S., 12 pp. Cont.-in-part of U.S. Ser. No. 166,144. CODEN: USXXAM

DT Patent

LA English

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
				<del>-</del>	
PI	US 5463116	Α	19951031	US 1994-190460	19940202
	US 5488150	Α	19960130	US 1993-166144	19931214
	CA 2114678	AA	19950802	CA 1994-2114678	19940201
	CA 2114678	C	19990427		
PRAI	JP 1991-189696	A	19910730		
	JP 1991-199453	A	19910808		
	US 1992-921224	B1	19920729		
	US 1993-166144	A2	19931214		
ΔR	Stable crystals of	N_ (tran	g - 4 - i gonzoni	d creal about least a sell a	1 7 7

AB Stable crystals of N-(trans-4-isopropylcyclohexylcarbonyl)-D-phenylalanine for pharmaceutical formulation may be produced by treating this compound with a solvent at a temperature of at least 10° and forming crystals in the solvent at a temperature of at least 10°. For example, crystals may be formed by crystallization out of solution, or may be formed from solid particles

of the compound suspended in a solvent. Crystals formed in this way have different m.p., IR spectrum and X-ray diffraction patterns from previously known forms of the compound and have enhanced processability, e.g., stability to grinding.

IT 105816-04-4

RL: PEP (Physical, engineering or chemical process); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(crystallization of (isopropylcyclohexylcarbonyl)phenylalanine for enhanced stability to grinding)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

IT 173653-89-9

RL: PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(crystallization of (isopropylcyclohexylcarbonyl)phenylalanine for enhanced stability to grinding)

RN 173653-89-9 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]-, hydrate (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

#### ●x H<sub>2</sub>O

L15 ANSWER 35 OF 35 CAPLUS COPYRIGHT 2004 ACS on STN

AN 1993:261002 CAPLUS

DN 118:261002

TI Stable **crystals** of N-(trans-4-isopropylcyclohexylcarbonyl)-D-phenylalanine

IN Sumikawa, Michito; Koguchi, Yoshihito; Ohgane, Takao; Irie, Yasuo; Takahashi, Satoji

PA Ajinomoto Co., Inc., Japan

SO Eur. Pat. Appl., 14 pp. CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 2

I I III	CIVI				
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
			- <b></b>		
PΙ	EP 526171	A2	19930203	EP 1992-306895	19920729
	EP 526171	A3	19930505		23320,23

EP	526171	B1	19970305		
	R: AT, CH,	DE, DK,	ES, FR, GB,	IT, LI, LU, NL, SE	
JP	05208943	A2	19930820	JP 1992-202686	19920729
JP	2508949	B2	19960619		
AT	149483	E	19970315	AT 1992-306895	19920729
ES	2100291	Т3	19970616	ES 1992-306895	19920729
CA	2114678	- AA	19950802	CA 1994-2114678	19940201
CA	2114678	С	19990427		
PRAI JP	1991-189696	A	19910730		
JР	1991-199453	A	19910808		
		_	_		

AB Stable H-type crystals of N-(trans-4-isopropylcyclohexylcarbonyl)-D-phenylalanine (I) are obtained by treating I with a solvent, at >10°. A solution of 5 g I in 20 mL acetone was added to a stirred mixture of 40 mL acetone and 60 mL water, at 25° to precipitate H-type crystals. The crystals have different m.p., IR spectrum and x-ray diffraction patterns from known forms of I and are not converted to other forms when ground.

IT 105816-04-4P

RL: PREP (Preparation)

(crystals, stable, preparation of)

RN 105816-04-4 CAPLUS

CN D-Phenylalanine, N-[[trans-4-(1-methylethyl)cyclohexyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

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FILE 'WPIDS' ENTERED AT 12:16:34 ON 20 SEP 2004 COPYRIGHT (C) 2004 THOMSON DERWENT

FILE LAST UPDATED: 15 SEP 2004 <20040915/UP>
MOST RECENT DERWENT UPDATE: 200459 <200459/DW>

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searched by Alex Waclawiw Page 43

FIRST VIEW - FILE WPIFV.

searched by Alex Waclawiw Page 44

FOR FURTHER DETAILS: http://www.thomsonderwent.com/dwpifv <<< >>> NEW DISPLAY FORMAT HITSTR ADDED ALLOWING DISPLAY OF HIT STRUCTURES WITHIN THE BIBLIOGRAPHIC DOCUMENT <<< => d que 14,3 ≈ L43 NOT FOUND => d que 13 95 SEA FILE=WPIDS ABB=ON PLU=ON NATEGLINIDE OR STARLIX OR L1FASTIC OR DJN 608 OR SENAGLINIDE OR STARSIS OR AY 4166 OR A 4166 OR SDZ DJN 608 362002 SEA FILE=WPIDS ABB=ON PLU=ON CRYS? L216 SEA FILE=WPIDS ABB=ON PLU=ON L1 AND L2 L3 $=> d \cdot wp 13 1-16$ COPYRIGHT 2004 THOMSON DERWENT on STN L3ANSWER 1 OF 16 WPIDS AN2004-594140 [57] WPIDS 2004-108803 [11]; 2004-180282 [17] CR DNC C2004-216153 New crystalline nateglinide form-U useful to reduce TIblood glucose level and to treat type-II diabetes. DC B05 FRENKEL, G; GOME, B; WIZEL, S IN(TEVA-N) TEVA PHARM IND LTD; (TEVA-N) TEVA PHARM USA INC PACYC **1**07 PIWO 2004067496 A1 20040812 (200457) \* EN RW: AT BE BG) BW CH CY CZ DE DK EA EE ES FI FR GB GH GM GR HU IE IT KE <del>ls lu m</del>c mw mz nl oa pt ro sd se si sk sl sz tr tz ug zm zw W: AE AG AL AM AT AU AZ BA BB BG BR BW BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE EG ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NA NI NO NZ OM PG PH PL PT RO RU SC SD SE SG SK SL SY TJ TM TN TR TT TZ UA UG UZ VC VN YU ZA ZM ZW ADT WO 2004067496 A1 WO 2004-US839 20040113 PRAI US 2003-746697 20031224; US 2003-442109P 20030123; US 2003-449791P 20030224; US 2003-479016P 20030616; 20030718; WO 2003-US22375 US 2003-622905 20030718; US 2003-693166 20031023 ABWO2004067496 A UPAB: 20040907 NOVELTY - Crystalline nateglinide form-U (I) substantially free of a peak at 3.8 plus or minus 0.2 theta is new. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for (1) the preparation of (I); and (2) a process for purifying (I). ACTIVITY - Antidiabetic. No biological data given. MECHANISM OF ACTION - None given. USE - (I) is useful to lower blood sugar level and to treat type II diabetes (claimed). ADVANTAGE - (I) has improved pharmaceutical characteristics such as targeted release profile. Dwg.0/69 TECH UPTX: 20040907 TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preparation (Claimed): Preparation

of (I) comprises (a) either preparing a solution of nateglinide in ethyl acetate at about 40-45 degrees C and adding in any order a 5-12C aliphatic hydrocarbon having about 5 degrees C as an anti-solvent to precipitate nateglinide; or (b) preparing a solution of nateglinide in ethylacetate, seeding the solution with nateglinide crystals and recovering the crystalline form as a precipitate; or (c) preparing a container holding a solution of nateglinide in ethyl acetate, adding 5-12C hydrocarbon to the container holding the solution and recovering the crystalline form as a precipitate; or (d) preparing a solution of nateglinide in a mixture of water and ethyl acetate, combining the solution with an anti-solvent and recovering the crystalline nateglinide as a precipitate. Purification of (I) comprises crystallizing the crystalline nateglinide from a solution in the presence of water resulting in the crystalline form being 99% pure as area percentage high performance liquid chromatography. Preferred Components: (I) has a XRPD pattern with peaks at about 4.7, 7.4, 13.8 and 17.0+/-0.2 and a FTIR spectrum with peaks at about 3350, 1701, 1646 and 1291 cm-1. Preferred Process: The antisolvent is heptane. The volume of ethyl acetate is 3-11 (preferably 4-6) ml/g compared to weight of the nateglinide. The solution is seeded with the same crystalline form. The preparation further comprises cooling before or after seeding and seeding before precipitation. (I) obtained is free of other crystalline forms by weight. The hydrocarbon is a 5-8C hydrocarbon (preferably heptane) and is added in such a manner to avoid precipitation upon addition. The precipitation is carried out in the presence of water. The recovering of (I) is by filtering the precipitate. The preparation also comprises preparing a solution of nateglinide in ethyl acetate at 25-50 degrees C at an ethyl acetate/ nateglinide ratio of about 3 -1 ml/g, seeding the solution with the same crystalline nateglinide at about 10-35 degrees C, stirring the seeded solution, cooling the seeded solution at a rate of about 1-10 degrees C per hour to a temperature of about (-)10-10 degrees C, filtering the crystalline nateglinide as a precipitate and drying the precipitate. The ethyl acetate is mixed with water at about 2-10% water as percentage of milliliters of water to grams of nateglinide. The anti-solvent is a 5-12C hydrocarbon (preferably heptane). In the purification process the rest of the solution is comprised of ethyl acetate. ANSWER 2 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN

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L3
     2004-26919 [25]
AN
                        WPIDS
DNC
     C2004-104807
     New crystalline form of nateglinide useful to treat
     diabetes and to stimulate insulin secretion from pancreas.
DC
     B05
     KADABOINA, R; POLAVARAPU, S; REGURI, B R
IN
PA
     (REDD-N) REDDY'S LAB LTD
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A1 20040311 (200425)\* EN

29

RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GH GM GR HU IE IT KE LS LU MC MW MZ NL OA PT RO SD SE SI SK SL SZ TR TZ UG ZM ZW W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NI NO NZ OM PG PH

CYC

PI

105

WO 2004020396

PL PIORO RU SC SD SE SG SK SL SY TJ TM TN TR TT TZ UA UG US UZ VC VN YU ZA ZM ZW US/2004077725\ A1 20040422 (200428) ADT WO 2004020396 A1 WO 2003-US26880 20030827; US 2004077725 A1 US 2003-649380 200330827 PRAI IN 2002-CH631/ 20020828 WO2004020396 A UPAB: 20040525 NOVELTY - Crystalline form X of nateglinide (I) is new. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for (1) a composition (B) comprising nateglinide as a solid, where at least 80% by weight of the solid is (I); and (2) preparation of (I). ACTIVITY - Antidiabetic. MECHANISM OF ACTION - None given in the source material. USE - (I) is useful to treat diabetes and also stimulates the secretion of insulin from pancreas. Dwg.0/2TECH UPTX: 20040418 TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preparation (Claimed): Preparation of (I) comprises: (a) providing a solution of nateglinide in an aromatic hydrocarbon solvent; (b) cooling the solution until a precipitate is formed; and (c) isolating (I). Preferred Process: The starting nateglinide is crystalline form H and/or B. The hydrocarbon solvent is benzene, ethylbenzene and toluene (preferably xylene or ortho-xylene). The process further comprises heating the mixture of the starting material and hydrocarbon solvent at 40 degrees C - 130 degrees C (preferably 50 degrees C - 70 degrees C), drying the isolated precipitate and filtering the nateglinide solution before cooling. Cooling is carried out at 25 degrees C - 35 degreesC TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Components: (I) exhibits an X-ray diffraction pattern expressed in terms of 2 theta angles, that includes 5 or more peaks of 3.95+/-0.09, 4.89+/-0.09, 5.18+/-0.09, 6.78 + /-0.09, 7.79 + /-0.09, 10.32 + /-0.09, 13.51 + /-0.09, 14.00 + /-0.09, 16.98 + (-0.09, 17.94 + (-0.09, 18.85 + (-0.09, 19.17 + (-0.09, 20.32 + (-0.09, 16.98 + (-0.09, 19.17 + (-0.21.12+/-0.09, 22.52+/-0.09, 23.76+/-0.09, 24.46+/-0.09, 27.36+/-0.09, 28.17 + (-0.09, 30.88 + (-0.09, 31.25 + (-0.09, 32.61 + (-0.09, and 41.65 + (-0.09))degrees (particularly 3.95+/-0.09, 14.00+/-0.09, 16.98+/-0.09) against Lin (counts per second) and 2 theta angles. The X-ray diffraction pattern further includes 3.952, 14.039, 16.98, 20.325, 21.120, 17.942, 6.776, 13.515 and 18.853 degrees. (I) also exhibits an infrared absorption spectrum with absorption bands at about 3353 cm-1, 2937cm-1, 2868 cm-1, 1743 cm-1, 1646 cm-1, 1597 cm-1, 1445 cm-1, 1208 cm-1, 1190 cm-1, 1110 cm-1, 697 cm-1 and 607 cm-1 against T% and cm-1. Preferred Composition: (B) comprises at least 90% (preferably 99%) by weight of (I) and at least 1% (preferably 5%) solid nateglinide is not its crystalline form. The solid nateglinide is substantially free of its crystalline forms B and H. ANSWER 3 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN L32004-180282 [17] ANWPIDS 2004-108803 [11]; 2004-594140 [57] CR DNC C2004-071244 New crystalline polymorphic forms of nateglinide

useful for lowering the blood sugar level.

TI

DC

B05

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DOLITZKY, B; GOME, B; GOZLAN, Y; SHAPIOR, E; YAHALOMI, R; SHAPIRO, E
 IN
      (TEVA-N) TEVA PHARM IND LTD; (DOLI-I) DOLITZKY B; (GOME-I) GOME B;
PA
      (GOZL-I) GOZLAN Y; (SHAP-I) SHAPIRO E; (YAHA-I) YAHALOMI R; (TEVA-N) TEVA
     PHARM USA INC
CYC
     105
     WØ 2004009532
PI
                     A1 20040129 (200417) * EN 130
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            LU MC MW MZ NL OA PT RO SD SE SI SK SL SZ TR TZ UG ZM ZW
         W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK
            DM DZ EC EE ES FI GB GD GE GH GM. HR HU ID IL IN IS JP KE KG KP KR
            KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NI NO NZ OM PG PH
            PL PT RO RU SC SD SE SG SK SL SY TJ TM TN TR TT TZ UA UG US UZ VC
            VN YU ZA ZM ZW
     US 2004116526 A1 20040617 (200440)
     AU 2003253971 A1 20040209 (200450)
     US 2004152782 A1 20040805 (200452)
ADT WO 2004009532 A1 WO 2003-US22375 20030718; US 2004116526 A1 Provisional US
     2002-396904P 20020718, Provisional US 2002-413622P 20020925, Provisional
     US 2002-414199P 20020926, Provisional US 2002-423750P 20021105,
     Provisional US 2002-432093P 20021210, Provisional US 2002-432962P
     20021212, Provisional US 2003-442109P 20030123, Provisional US
     2003-449791P 20030224, Provisional US 2003-479016P 20030616, US
     2003-623237 20030718; AU 2003253971 A1 AU 2003-253971 20030718; US
     2004152782 A1 Provisional US 2002-393495P 20020703, Provisional US
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     US 2002-414199P 20020926, Provisional US 2002-423750P 20021105,
     Provisional US 2002-432093P 20021210, Provisional US 2002-432962P
     20021212, Provisional US 2003-442109P 20030123, Provisional US
     2003-449791P 20030224, US 2003-614266 20030703
FDT AU 2003253971 A1 Based on WO 2004009532
PRAI US 2003-614266
                          20030703; US 2002-396904P
                                                         20020718;
     US 2002-413622P
                          20020925; US 2002-414199P
                                                         20020926;
     US 2002-423750P
                          20021105; US 2002-432093P
                                                         20021210;
     US 2002-432962P
                          20021212; US 2003-442109P
                                                         20030123;
     US 2003-449791P
                          20030224; US 2003-479016P
                                                         20030616;
     US 2003-623237
                          20030718; US 2002-393495P
                                                         20020703
AB
     WO2004009532 A UPAB: 20040907
     NOVELTY - 26 Crystalline nateglinide forms as
     characterized by XRPD patterns, DSC thermograms and FTIR spectra, fully
     described in the specification, are new.
          DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the
     preparation of the crystalline forms of nateglinide.
          ACTIVITY - Antidiabetic.
          No test details for antidiabetic activity are given.
          MECHANISM OF ACTION - None given.
         USE - The pharmaceutical formulation comprising crystalline
     nateglinide form of A, C, D, F, G, I, J, K, M, NO, Q, T, V, Y,
     gamma, epsilon, theta or omega is useful to lower the blood sugar level
     (claimed).
         ADVANTAGE - The new polymorphic forms of nateglinide
     provides a new opportunity to improve the performance characteristics of a
     pharmaceutical product.
    Dwg.0/64
TECH
                   UPTX: 20040310
     TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preparation (Claimed): Preparation
    of nateglinide form B comprises preparing a suspension of
    nateglinide in a 5-6C hydrocarbon, adding a solvent of an alcohol,
    ester and/or ketone to the suspension to obtain a solution,
    crystallizing nateglinide form B from the solution in
    the absence of stirring and recovering the nateglinide form B.
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The crystallization is carried out by seeding and cooling at higher than 15 degrees C. Preparation of other crystal forms

comprises heating one crystal form to obtain another or

extracting the crystal form from a solvent.

Preferred Reagents: The hydrocarbon is heptane, hexane, toluene and xylene. The solvent is methanol, ethanol, isopropanol, n-butanol, n-propanol, acetone or ethyl acetate. ANSWER 4 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN L32004-108803 [11] WPIDS AN2004-180282 [17]; 2004-594140 [57] CR DNC C2004-044538 Preparation of trans-4-isopropylcyclohexane acid chloride as intermediate  $\mathbf{T}\mathbf{I}$ in preparing nateglinide comprises reaction between thionyl chloride and acid chloride in the presence of organic amide. DCB05 DOLITZKY, B; GOME, B; GOZLAN, Y; SHAPIOR, E; YAHALOMI, R; SHAPIRO, E IN(TEVA-N) TEVA PHARM IND LTD; (TEVA-N) TEVA PHARM USA INC PACYC 105 W2 2004005240 PIA1 20040115 (200411)\* EN 31 RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GH GM GR HU IE IT KE LS LU MC/ MW MZ NL OA PT RO SD SE SI SK SL SZ TR TZ UG ZM ZW W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NI NO NZ OM PG PH PL PT RO RU SC SD SE SG SK SL SY TJ TM TN TR TT TZ UA UG US UZ VC VN YU ZA ZM ZW A1 20040209 (200450) AU 2003253971 Al 20040123 (200459) AU 2003256454 ADT WO 2004005240 A1 WO 2003-US21238 20030703; AU 2003253971 A1 AU 2003-253971 20030718; AU 2003256454 A1 AU 2003-256454 20030703 FDT AU 2003253971 Al Based on WO 2004009532; AU 2003256454 Al Based on WO 2004005240 PRAI US 2003-479016P 20030616; US 2002-393495P 20020703; US 2002-396904P 20020718; US 2002-413622P 20020925; US 2002-414199P 20020926; US 2002-423750P 20021105; US 2002-432093P 20021210; US 2002-432962P 20021212; US 2003-442109P 20030123; US 2003-449791P 20030224; US 2003-614266 20030703 WO2004005240 A UPAB: 20040915 ABNOVELTY - Preparing trans-4-isopropylcyclohexane acid chloride comprises combining trans-4-isopropylcyclohexane carboxylic acid with thionyl chloride in the presence of a 1-6C organic amide to obtain trans-4-isopropylcyclohexane acid chloride free of its corresponding cis isomer; and recovering the trans-4-isopropylcyclohexane acid chloride. DETAILED DESCRIPTION - An INDEPENDENT CLAIM is also included for a process for preparing nateglinide by combining trans-4-isopropylcyclohexane carboxylic acid with thionyl chloride in the presence of a 1-6C organic amide to obtain trans-4-isopropylcyclohexane acid chloride free of its corresponding cis isomer; converting the acid chloride to nateglinide; and recovering the nateglinide ACTIVITY - Antidiabetic. MECHANISM OF ACTION - None given.

USE - For preparing trans-4-isopropylcyclohexane acid chloride as an

ADVANTAGE - The cis-isomer is not formed nor detected in amounts of

intermediate in preparing nateglinide for the treatment of type

less than 0.05% even at elevated temperature (60-80 deg. C) in the

reaction between thionyl chloride and trans-isopropylcyclohexane

II diabetes.

carboxylic acid in the presence of an organic amide catalyst. Dwg.0/3TECH UPTX: 20040213 TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Process: The combining is carried out by adding 0.05-10 wt.% amide to 1-5 acid equivalents of thionyl chloride at 10-60 degrees C. The reaction mixture is then maintained for 1-5 hours. The nateglinide may be prepared by combining a solution of a tri-alkyl amine salt of D-phenylalanine with trans-4-isopropylcyclohexane acid chloride in a 1-7C amide to form nateglinide; and recovering the nateglinide. The nateglinide may be prepared in a two-phase system by preparing an aqueous solution of an alkaline earth or alkali metal salt of D-phenylalanine; combining the aqueous solution with a water-immiscible organic solvent containing trans-4-isopropylcyclohexane acid chloride to form an aqueous and an organic phase, wherein nateglinide forms through reaction between the D-phenylalanine and the trans-4isopropylcyclohexane acid chloride; and recovering the nateglinide . Preparing nateglinide Form Z comprises preparing an aqueous solution of an alkali metal or an alkali earth metal salt of nateglinide; acidifying the solution to precipitate nateglinide; and recovering the nateglinide Form Z. The aqueous solution contains water free of a co-solvent. Preparing nateglinide further comprises crystallizing /recrystallizing. Preferred Composition: The weight ratio of the cis isomer to the trans isomer is less than 0.03%. Preferred Compounds: The combining is carried out in a solvent, preferably aromatic or saturated hydrocarbon, ester or ether. The tri-alkyl amine is triethyl amine. The organic amide is N, N-dimethylacetamide, N-methylpyrrolidone or preferably N, N-dimethylformamide. The water-immiscible organic solvent is 5-12C hydrocarbon, preferably toluene, heptane or ethyl acetate. ANSWER 5 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN L32004-081844 [08] ANWPIDS DNC C2004-033612 New crystal form of N-(trans-4-isopropylcyclohexylcarbonyl)-Dphenylalanine useful for lowering blood glucose level. DCA96 B05 SUTTON, PA IN(NOVS) NOVARTIS AG; (NOVS) NOVARTIS PHARMA GMBH PACYC 90 WO 2003087038 PIA1 20031023 (200408) \* EN RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GR HU IE IT LU MC NL PT RO SE SI SK TR W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LT LU LV MA MD MK MN MX NI NO NZ OM PH PL PT RO RU SC SE SG SK TJ TM TN TR TT UA US UZ VC VN YU ZA ZW Al 20031027 (200436) AU 2003242520 ADT WO 2003087038 A1 WO 2003-EP3864 20030414; AU 2003242520 A1 AU 2003-242520 20030414 FDT AU 2003242520 Al Based on WO 2003087038 PRAI US 2002-372625P 20020415 AB WO2003087038 A UPAB: 20040202 NOVELTY - A crystal form of N-(trans-4isopropylcyclohexylcarbonyl) -D-phenylalanine (nateglinide) having melting point of 108 deg. C, or its solvate is new.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for the

production of R'-type crystal form of nateglinide

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involving:
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- (a) dissolving nateglinide in any of its forms in a solvent (S1) in which nateglinide is readily soluble at an ambient temperature to form a solution;
- (b) treating the solution with another solvent (S2) which is miscible with (S1) and in which nateglinide is poorly soluble to induce precipitation of R'-type crystals of nateglinide; and
- (c) isolating and drying the precipitate crystal form of nateglinide.

ACTIVITY - Antidiabetic.

MECHANISM OF ACTION - None given.

USE - For lowering blood glucose level in human.

ADVANTAGE - The nateglinide in any of its form, such as hydrates, methanolates, ethanolates and acetonates can be used for the production of R'-type crystal. Dwq.0/2

TECH

UPTX: 20040202

TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Method: The crystal form of nateglinide is induced by stirring, cooling or adding seed crystals of nateglinide. The ambient temperature is from room temperature to the boiling point of the solvent (preferably room temperature). The crystal form of nateglinide is dried under atmospheric or reduced pressure (preferably reduced pressure) at room temperature to 70 degrees Celsius (preferably room temperature to 50 degrees Celsius). Preferred Components: (S1) is a mixture of ethanol (50 vol.%) and toluene. (S2) is water containing hydroxypropylmethylcellulose (1%). The ratio of (S1) and (S2) is 1:7 vol.

ANSWER 6 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN L3

2003-853914 [79] ANWPIDS

DNC C2003-240851

New crystalline nateglinide forms A, M and P are TIantiglycemic agents and antidiabetic agents.

B05 DC

KOGUCHI, Y; NAKAO, T; SUMIKAWA, M IN

(AJIN) AJINOMOTO CO INC PA

CYC 103

A1 20031023 (200379)\* JA WO 2003087039

RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GH GM GR HU IE IT KE LS LU MC MW MZ NL OA PT RO SD SE SI SK SL SZ TR TZ UG ZM ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NI NO NZ OM PH PL PT RO RU SC SD SE SG SK SL TJ TM TN TR TT TZ UA UG US UZ VC VN YU ZA ZM ZW

AU 2003236243 A1 20031027 (200436)

WO 2003087039 A1 WO 2003-JP4686 20030414; AU 2003236243 A1 AU 2003-236243 ADT20030414

AU 2003236243 A1 Based on WO 2003087039 FDT

PRAI JP 2002-111963 20020415

WO2003087039 A UPAB: 20031208 AB

> NOVELTY - Crystalline nateglinide forms A, M and P are new.

> DETAILED DESCRIPTION - Crystalline nateglinide of formula (I) forms A, M and P are new.

USE - Nateglinide is an antiglycemic agent and antidiabetic agent.

ADVANTAGE - Have improve stability and solubility. Dwg.0/3

TECH

UPTX: 20031208

TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Crystalline Forms: Crystalline forms have the following powder X-ray diffraction peaks (2 theta) Form A 4.4, 5.2, 15.7 and 18.5 degrees; form M 6.0, 14.2, 15.2 and 18.8 degrees and form P 4.8, 5.3 14.3 and 15.2 degrees.

Preparation: Preparation of crystalline A form e.g. comprises: (1) dissolving nateglinide in a solvent having high solubility for nateglinide and adding a solvent with poor solubility for nateglinide or dissolving nateglinide in a mixture of solvents having high and poor solubility for nateglinide (preferably ethanol and water);

- (2) cooling the mixture (preferably to 10 degrees C) to precipitate crystals; and
- (3) filtering and drying at 30-80 (preferably 40-60) degrees C. Preparation of crystalline form A and P form comprises e.g. heating crystalline form B at 60 degrees C or more (preferably 80 degrees C or more). Crystalline form M is prepared e.g. by heating crystalline form B at 40-100 (preferably 50-70) degrees C and 60-95 (preferably 70-90)% relative humidity.
- L3 ANSWER 7 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN

AN 2003-748369 [70] WPIDS

DNC C2003-205231

New salt of **nateglinide** useful for treating, e.g. diabetes, cardiovascular or related diseases, e.g. hyperglycemia, hyperlipidaemia, obesity, diabetes retinopathy, diabetic neuropathy, glomerulosclerosis or stroke.

DC B05

IN DE LA CRUZ, M; PARKER, D J; SUTTON, P A; VIVILECCHIA, R V

PA (NOVS) NOVARTIS AG; (NOVS) NOVARTIS PHARMA GMBH

CYC 90

PI WO 2003076393 A1 20030918 (200370) \* EN 23

RW: AT BE BG CH CY CZ DE DK EA EE ES FI FR GB GR HU IE IT LU MC NL PT RO SE SI SK TR

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LT LU LV MA MD MK MN MX NI NO NZ OM PH PL PT RO RU SC SE SG SK TJ TM TN TR TT UA US UZ VC VN YU ZA ZW

AU 2003214112 A1 20030922 (200431)

ADT WO 2003076393 A1 WO 2003-EP2447 20030310; AU 2003214112 A1 AU 2003-214112 20030310

FDT AU 2003214112 Al Based on WO 2003076393

PRAI US 2002-363178P 20020311

AB W02003076393 A UPAB: 20031030

NOVELTY - A salt of **nateglinide** (I) having a melting point of 50-300 deg. C is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for:

(1) A composition comprising (I); and

(2) A method for the treatment of diabetes, cardiovascular disease or related conditions, comprising administration of (I).

ACTIVITY - Antidiabetic; Antilipemic; Anorectic; Ophthalmological; Neuroprotective; Nephrotropic; Vasotropic; Antiulcer; Antiinflammatory; Cardiant; Hypotensive; Antianginal; Cerebroprotective; Dermatological; Antiarthritic; Osteopathic; Vasotropic; Cardiovascular-Gen.

Test details are described, but no results given.

MECHANISM OF ACTION - None given.

USE - (I) is used for treating diabetes, cardiovascular or related diseases, e.g. hyperglycemia, hyperinsulinaemia, hyperlipidaemia, insulin resistance, impaired glucose metabolism, obesity, diabetes retinopathy,

macular degeneration, cataracts, diabetic neuropathy, glomerulosclerosis, erectile dysfunction, premenstrual syndrome, vascular restenosis, ulcerative colitis, coronary heart disease, hypertension, angina pectoris, myocardial infarction, stroke, skin and connective tissue disorder, foot ulcerations, metabolic acidosis, arthritis, osteoporosis, polycystic ovary syndrome or impaired glucose tolerance (all claimed).

ADVANTAGE - The salt of nateglinide has a higher degree of dissociation in water, increased biological availability of the salts, salt hydrates, or salt anions in the case of solid dosage forms. For different relative humidities at room temperature, the salts shows (with the exception of potassium and a calcium salt) practically no water absorption or water loss over a wide range of humidities and for periods of few hours, e.g. four hours. The melting point of the salts will not be changed by storing under different relative humidities, except for the melting point of those salts that are hygroscopic or moderately hygroscopic. (I) has a water solubility of at least 0.18 (preferably at least 0.4, especially 40) mg/ml.

TECH

UPTX: 20031030

TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Compound: (I) is confirmed by X-ray powder diffraction (XRPD) pattern and is present in an amorphous and/or crystalline form.

The ratio of nateglinide anion to cation is 1:1 or 2:1.

The salt loses 0.1-14 (preferably 0.1-0.9)% of its mass on heating. (I) has a bulk density of 0.1-0.6 q/cm3.

Preferred Cation: In (I), the cation is a sodium ion (Na+), potassium ion (K+), calcium ion (Ca2+), magnesium ion (Mg2+) or the protonated form of tris(hydroxymethyl)-aminomethane or N-methyl-D-glucamine or lysine. Preferred Composition: The composition also comprises at least one of vitamins, nutrition supplements, active substances, nateglinide or repaglinide.

The active substance is an insulin sensitizer, insulin secretion enhancer, dipeptidyl peptidase IV inhibitor, ACE inhibitor or angiotensin II inhibitor.

L3 ANSWER 8 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN

AN 2003-111806 [10] WPIDS

DNC C2003-028518

New crystalline complex between either (D) or (L) enantiomers of natural amino acids and amorphous C-aryl glucoside compounds useful for treating e.g. diabetes.

DC B03

IN GOUGOUTAS, J Z

PA (GOUG-I) GOUGOUTAS J Z; (BRIM) BRISTOL-MYERS SQUIBB CO

CYC 101

PI WO 2002083066 A2 20021024 (200310) \* EN 80

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZM ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ OM PH PL PT RO RU SD SE SG SI SK SL TJ TM TN TR TT TZ UA UG US UZ VN YU ZA ZM ZW

US 2003064935 A1 20030403 (200325)

EP 1385856 A2 20040204 (200410) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR

AU 2002254567 Al 20021028 (200433)

US 6774112 B2 20040810 (200453)

ADT WO 2002083066 A2 WO 2002-US11066 20020408; US 2003064935 A1 Provisional US

2001-283097P 20010411, US 2002-117914 20020408; EP 1385856 A2 EP 2002-723801 20020408, WO 2002-US11066 20020408; AU 2002254567 A1 AU 2002-254567 20020408; US 6774112 B2 Provisional US 2001-283097P 20010411, US 2002-117914 20020408

FDT EP 1385856 A2 Based on WO 2002083066; AU 2002254567 A1 Based on WO 2002083066

PRAI US 2001-283097P 20010411; US 2002-117914 20020408

AB WO 200283066 A UPAB: 20030211

NOVELTY - A crystalline complex between either (D) or (L) enantiomers of natural amino acid and amorphous C-aryl glucoside compound is new.

DETAILED DESCRIPTION - **Crystalline** complexes between either (D) or (L) enantiomers of natural amino acids and compound of formula (I) are new.

R1, R2 and R2a = H, OH, OR5, alkyl, -OCHF2, -OCF3, -SR5a or halo;
R3 and R4 = H, OH, OR5b, (cyclo)alkyl, CF3, -OCHF2, -OCF3, halogen,
-CONR6R6a, -CO2R5c, -CO2H, -COR6b, -CH(OH)R6c, -CH(OR5d)R6d, -CN,
-NHCOR5e, -NHSO2R5f, -NHSO2Aryl, -SR5g, -SOR5h, SO2R5i or 5 - 7-membered heterocycle (containing 1 - 4 heteroatoms of N, O, S, SO and/or SO2);

R3+R4 and NR6+R6a = annelated 5 - 7-membered carbocycle or heterocycle (both containing 1 - 4 heteroatoms of N, O, S, SO and/or SO2));

R5 and R5a - R5i = alkyl;

R6 and R6a - R6d = H, alkyl, (alkyl) aryl or cycloalkyl.

INDEPENDENT CLAIMS are included for the following:

- (1) A pharmaceutical combination (A1) comprising complex of either the (D) or (L) enantiomer of natural amino acids with (I) and a component (G1) selected from an antidiabetic agent (G) other than an SGLT2 inhibitor, an agent for treating the complications of diabetes, an anti-obesity agent, an antihypertensive agent, an antiplatelet agent, an antiatherosclerotic agent and/or a lipid-lowering agent (preferably G); and
- (2) Treating type II diabetes involving administering the complex of (I) alone or in combination with another antidiabetic agent, an agent for treating the complications of diabetes, an anti-obesity agent, an antihypertensive agent, an antiplatelet agent, an antiatherosclerotic agent and/or a hypolipidemic agent.

ACTIVITY - Antidiabetic; Ophthalmological; Neuroprotective; Vulnerary; Anorectic; Antiarteriosclerotic; Hypotensive; Nephrotropic. MECHANISM OF ACTION - Inhibitors of sodium dependent glucose transporters.

USE - Compound (I) is used for treating or delaying the progression or onset of diabetes, diabetic retinopathy, diabetic neuropathy, diabetic nephropathy, delayed wound healing, insulin resistance, hyperglycemia, hyperinsulinemia, elevated blood levels of fatty acids or glycerol, hyperlipidemia, obesity, hypertriglyceridemia, Syndrome X, diabetic complications, atherosclerosis or hypertension or for increasing high density lipoprotein levels and for treating type II diabetes (claimed).

ADVANTAGE - The complex normalizes the plasma glucose by enhancing the excretion of glucose in the urine, thus improves insulin sensitivity and delays the development of diabetic complications. Dwg.0/0

TECH UPTX: 20030211

TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preparation: The complex is prepared by:

- (1) Dissolving (I) in a water miscible solvent that is heated to 50 80 degrees C;
- (2) Transferring the solution rapidly to a 50 80 degrees C aqueous or alcoholic solution containing either one or two equivalents of either the (D) or (L) enantiomer of a natural amino acid; and

(3) Upon slowly cooling, isolating the crystals of the desired complex form by filtration. Preferred Compound: The compound is a compound of formula (Ia). R'1 = H, alkoxy, halogen or lower alkyl; R'4 = lower alkyl, R5aO, -OCHF2, SR5e, S(O)R5e or S(O)2R5e.Preferred Complex: The complex comprises L-phenylalanine:1-(-(3-(4difluoromethoxybenzyl)-4-methylphenyl)-beta-D-glucopyranoside in a ratio of 1:1, L-Phenylalanine:1-(-(3-(4-methylthiobenzyl)-4-methylphenyl)-beta-Dglucopyranoside in a ratio 1:1, L-phenylalanine:1-(-(3-(4-ethylbenzyl)phenyl)-beta-D-glucopyranoside in a ratio of 1:1, L-phenylalanine:1-(-(3-(4-ethylbenzyl)-phenyl)-beta-D-glucopyranoside in a ratio of 2:1, L-proline: 1-(-(3-(4-ethylbenzyl)-phenyl)-beta-D-glucopyranoside in a ratio of 2:1, L-proline:1-(-(3-(4-ethylbenzyl)-phenyl)-beta-D-glucopyranoside in a ratio of 1:1, L-proline:1-(-(3-(4-methylthiobenzyl)-4-methylphenyl)-beta-D-glucopyranoside in a ratio of 1:1 or D-phenylalanine:1-(-(3-(4methylthiobenzyl) -4-methylphenyl) -beta-D-glucopyranoside in a ratio of 1:1.

TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Combination: A weight ratio of the complex of either the (D) or (L) enantiomer to (G) or to the lipid-lowering agent is 0.01 - 300:1. Preferred Components: (G) is at least one of a biguanide, a sulfonyl urea, a glucosidase inhibitor, a PPAR gamma agonist, a PPAR alpha/gamma dual agonist, an aP2 inhibitor, a DP4 inhibitor, n insulin sensitizer, a glucagons-like peptide-1 (GLP-1), insulin, a meglitinide, a PTP1B inhibitor, a glycogen phosphorylase inhibitor and/or a glucose-6-phosphatase inhibitor, preferably at least one of metformin, glyburide, glimepride, glipyride, glipizide, chlorpropamide, gliclazide, acarbose, miglitol, pioglitazone, troglitazone, rosiglitazone, insulin, G1-262570, isaglitazone, JTT-501, NN-2344, L895645, YM-440, R-119702, AJ9677, repaglinide, nateglinide, KAD1129, AR-HO39242, GW-409544, KRP297, AC2993, LY315902 and/or NVP-DPPD-728A. The anti-obesity agent is a beta 3 adrenergic agonist, a lipase inhibitor, a serotonin and dopamine reuptake inhibitor, a thyroid receptor beta compound and/or an anorectic agent (preferably orlistat, ATL-962, AJ9677, L750355, CP331648, sibutramine, topiramate, axokine, dexamphetamine, phentermine, phenylpropanolamine and/or mazindol). The lipid lowering agent is an MTP inhibitor, an HMG CoA reductase inhibitor, a squalane synthetase inhibitor, a fibric acid derivative, an upregulator of LDL receptor activity, a lipoxygenase inhibitor or an ACAT inhibitor (preferably pravastatin, lovastatin, simvastatin, atorvastatin, cerivastatin, fluvastatin, nisvastatin, visastatin, atavastatin, rosuvastatin, fenofibrate, gemfibrozil, clofibrate, avasimbe, TS-962, MD-700 and/or LY295427).

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L3 ANSWER 9 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN
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AN 2002-713487 [77] WPIDS

DNC C2002~202321

Combination used for treating e.g. hypertension, obesity, diabetic neuropathy and arthritis comprises nateglinide or repaglinide and additional antidiabetic compound e.g. insulin.

DC B02 B05

IN VILLHAUER, E B

PA (NOVS) NOVARTIS AG; (NOVS) NOVARTIS PHARMA GMBH; (VILL-I) VILLHAUER E B; (NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH

CYC 88 PI WO

WO 2002072146 A2 20020919 (200277)\* EN 30

RW: AT BE CH CY DE DK EA ES FI FR GB GR IE IT LU MC NL PT SE TR

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK

DM DZ EC EE ES FI GB GD GE GH HR HU ID IL IN IS JP KE KG KP KR KZ

LC LK LT LU LV MA MD MK MN MX NO NZ OM PH PL PT RO RU SE SG SI SK TJ TM TN TR TT UA US UZ VN YU ZA ZW

EP 1385549 A2 20040204 (200410) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR

AU 2002254940 Al 20020924 (200433)

US 2004143015 A1 20040722 (200449)

ADT WO 2002072146 A2 WO 2002-EP2665 20020311; EP 1385549 A2 EP 2002-724221 20020311, WO 2002-EP2665 20020311; AU 2002254940 A1 AU 2002-254940 20020311; US 2004143015 A1 WO 2002-EP2665 20020311, US 2003-471253 20030910

FDT EP 1385549 A2 Based on WO 2002072146; AU 2002254940 A1 Based on WO 2002072146

PRAI US 2001-275098P 20010312; US 2003-471253 20030910

AB WO 200272146 A UPAB: 20021129

NOVELTY - Combination comprises nateglinide or repaglinide, at least one additional antidiabetic compound and optionally at least one carrier.

DETAILED DESCRIPTION - Combination comprises nateglinide or repaglinide, at least one additional antidiabetic compound and optionally at least one carrier. The antidiabetic compound comprises insulin signaling pathway modulator, compounds influencing a dys-regulated hepatic glucose production, pyruvate dehydrogenase kinase (PDHK) inhibitor, inhibitors of gastric emptying, insulin, inhibitors of glycogen synthase kinase-3, retinoid X receptor (RXR) agonists, agonists of human beta -3 adrenergic receptor, agonists of uncoupling proteins (UCPs), non-glitazone type PPAR- gamma , dual PPAR- gamma /PPAAR- alpha agonists, antidiabetic vanadium containing compounds, incretin hormones, beta -cell imidazoline receptor antagonist, miglitol or alpha 2-adrenergic antagonists.

The active ingredients are contained in the free form or in the form of their salts.

An INDEPENDENT CLAIM is also included for a commercial package comprising the combination together with instructions for simultaneous, separate or sequential used in the prevention, delay of progression or treatment of metabolic disorders or for improving the bodily appearance.

ACTIVITY - Antidiabetic; Ophthalmological; Anorectic; Nephrotropic; Vasotropic; Gynecological; Antiinflammatory; Antiulcer; Cardiant; Hypotensive; Cerebroprotective; Dermatological; Antiarthritic; Osteopathic.

MECHANISM OF ACTION - Iinsulin signaling pathway modulator; Pyruvate dehydrogenase kinase inhibitor; Retinoid X receptor agonist; Glycogen synthase kinase-3 inhibitor; Human beta -3 adrenergic receptor; Uncoupling protein agonist; beta -cell imidazoline receptor antagonist; Miglitol antagonist; alpha 2-adrenergic antagonist; Non-glitazone type PPAR- gamma, dual PPAR- gamma /PPAAR- alpha agonist.

No biological tests or results are given in the source material.

USE - Used for the prevention, delay of progression or treatment of metabolic disorders and for cosmetic treatment to obtain body weight loss (all claimed). The combination is used for treating hyperglycemia, hyperinsulinaemia, hyperlipidaemia, insulin resistance, impaired glucose metabolism, obesity, diabetic retinopathy, macular degeneration, cataracts, diabetic nephropathy, glomerulosclerosis, diabetic neuropathy, erectile dysfunction, premenstrual syndrome, vascular restenosis, ulcerative colitis, coronary heart disease, hypertension, angina pectoris, myocardial infarction, stroke, skin and connective tissue disorders, foot ulceration, metabolic acidosis, arthritis, osteoporosis and impaired glucose tolerance.

ADVANTAGE - The combination results in a beneficial, especially a synergistic, therapeutic effect. The combination also provides efficacy, a broader variety of therapeutic treatment and beneficial effects on

diseases and conditions associated with diabetes, which includes less gain of weight, compared to a mono-therapy applying only one of the active ingredients of the combination. Dwg.0/0

TECH

UPTX: 20021129

TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Compounds: The combination comprises 3-(4-(2-(2,3-dihydrobenzo(1,4)thiazin-4-yl)-ethoxy)-phenyl)-2-ethoxy propionic acid as a dual PPAR-gamma/PPAR-alpha agonist. The combination also includes at least one active compound comprising glitazone, sulfonylurea derivative, metformin, acarbose and/or their salts. The combination is in the form of a combined preparation or a pharmaceutical composition. The neteglinide is present in the B-type or H-type crystal modification.

L3 ANSWER 10 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN

AN 2002-507933 [54] WPIDS

DNC C2002-144389

Process for producing nateglinide crystals useful for treating diabetes involves reacting trans-4-isopropylcyclohexylcarbonyl chloride with D-phenylalanine in ketone and water in presence of alkali.

DC B05

IN NISHI, S; TAKAHASHI, D; TAKAHASHI, S

PA (AJIN) AJINOMOTO CO INC; (AJIN) AJINOMOTO KK

CYC 98

PI WO 2002032854 A1 20020425 (200254) \* JA 15

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PH PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001094265 A 20020429 (200255)

EP 1334963 A1 20030813 (200355) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR

BR 2001014729 A 20031014 (200374)

KR 2003059203 A 20030707 (200377)

US 2004030182 A1 20040212 (200412)

JP 2002536038 X 20040226 (200416) MX 2003003484 A1 20030701 (200423)

CN 1481356 A 20040310 (200437)

ADT WO 2002032854 Al WO 2001-JP9069 20011016; AU 2001094265 A AU 2001-94265 20011016; EP 1334963 Al EP 2001-974875 20011016, WO 2001-JP9069 20011016; BR 2001014729 A BR 2001-14729 20011016, WO 2001-JP9069 20011016; KR 2003059203 A KR 2003-705388 20030417; US 2004030182 Al Cont of WO 2001-JP9069 20011016, US 2003-418105 20030418; JP 2002536038 X WO 2001-JP9069 20011016, JP 2002-536038 20011016; MX 2003003484 Al WO 2001-JP9069 20011016, MX 2003-3484 20030416; CN 1481356 A CN 2001-820658 20011016

FDT AU 2001094265 A Based on WO 2002032854; EP 1334963 Al Based on WO 2002032854; BR 2001014729 A Based on WO 2002032854; JP 2002536038 X Based on WO 2002032854; MX 2003003484 Al Based on WO 2002032854

PRAI JP 2000-317604 20001018

AB WO 200232854 A UPAB: 20020823

NOVELTY - A process for producing nateglinide crystals involves:

- (i) reacting trans-4-isopropylcyclohexylcarabonyl chloride with D-phenylalanine in a mixed solvent, consisting of a ketone and water in the presence of an alkali; and
  - (ii) adding an acid to the resulting reaction mixture and subjected

to **crystallization** while regulating the temperature and the ketone solvent concentration.

DETAILED DESCRIPTION - A process for producing nateglinide crystals involves:

- (i) reacting trans-4-isopropylcyclohexylcarbonyl chloride with D-phenylalanine in a mixed solvent, consisting of a ketone and water in the presence of an alkali; and
- (ii) adding an acid, providing an acidic condition to the resulting reaction mixture, containing **nateglinide** and subjected to **crystallization** while regulating the temperature (between 58 72 deg. C) and the ketone solvent concentration (between 9 to up to but not including 22 wt%).

USE - For producing nateglinide crystals, which can be used as an oral medicine for treating diabetes.

ADVANTAGE - The process is efficient even on an industrial production scale.

Dwg.0/0

TECH UPTX: 20020823

TECHNOLOGY FOCUS - CHEMICAL ENGINEERING - Preferred Process: The adjustment of the ketone solvent concentration is conducted by adding a ketone (preferably acetone) to the reaction mixture, which is an acylation reaction solution. The ketone is at a concentration of 12 - 16 wt% in the reaction system.

Preferred Crystal: The crystal of the nateglinide is a H-type crystal having a mean long diameter of 1 mm or more and a mean short diameter of 0.1 mm or more.

- L3 ANSWER 11 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN
- AN 2002-500188 [53] WPIDS

DNC C2002-141632

- Hydrophilic drug preparation comprises nateglinide B crystals and has contact angle to water surface of 111 degrees or less useful as an hypoglycemic agent.
- DC A96 B05
- IN MAKINO, C; NINOMIYA, N; YABUKI, A
- PA (AJIN) AJINOMOTO CO INC; (AJIN) AJINOMOTO KK

CYC 98

- PI WO 2002040010 A1 20020523 (200253) \* JA 26
  - RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW
  - W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PH PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001096000 A 20020527 (200261)

EP 1334721 A1 20030813 (200355) EN

- R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR
- KR 2003042028 A 20030527 (200361)
- BR 2001014897 A 20030812 (200367)
- US 2004029968 A1 20040212 (200412)
- JP 2002542384 X 20040603 (200436)
- CN 1482904 A 20040317 (200437)
- ADT WO 2002040010 A1 WO 2001-JP9292 20011023; AU 2001096000 A AU 2001-96000 20011023; EP 1334721 A1 EP 2001-976818 20011023, WO 2001-JP9292 20011023; KR 2003042028 A KR 2003-705635 20030423; BR 2001014897 A BR 2001-14897 20011023, WO 2001-JP9292 20011023; US 2004029968 A1 Cont of WO 2001-JP9292 20011023, US 2003-420886 20030423; JP 2002542384 X WO 2001-JP9292 20011023, JP 2002-542384 20011023; CN 1482904 A CN 2001-821218 20011023
- FDT AU 2001096000 A Based on WO 2002040010; EP 1334721 Al Based on WO

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2002040010; BR 2001014897 A Based on WO 2002040010; JP 2002542384 X Based
     on WO 2002040010
PRAI JP 2000-324374
                           20001024
AB
     WO 200240010 A UPAB: 20020820
     NOVELTY - Hydrophilic drug preparation comprises nateglinide B
     crystals and has a contact angle to the surface of water of 111
     deg. or less.
          ACTIVITY - Antidiabetic.
          MECHANISM OF ACTION - None given.
          USE - As a hydrophillic drug preparation for administering
     nateglinide B crystals useful as an hypoglycemic agent.
          ADVANTAGE - Have quick release with high elution properties and are
     easily produced.
     Dwq.0/0
TECH
                    UPTX: 20020820
     TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preparation: Preparation is film or
     sugar coated, has a contact angle to the surface of water of 100
     (preferably 90) degrees or less and contains a hydrophilic substance
     (preferably a hydrophillic polymer, surfactant, sugar, sugar alcohol or
     salt)
     ANSWER 12 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN
L3
     2002-462521 [49]
AN
                        WPIDS
     1999-204733 [17]; 2000-170837 [15]; 2001-432562 [46]; 2001-522427 [57];
CR
     2001-595790 [67]; 2002-082346 [11]; 2002-215543 [27]; 2002-215909 [27];
     2002-315576 [35]; 2002-328338 [36]; 2002-635742 [68]; 2002-666828 [71];
     2002-696871 [75]; 2003-015683 [01]; 2003-198106 [19]; 2003-238931 [23];
     2003-417948 [39]; 2003-627162 [59]; 2003-776923 [73]
     N2002-364678
                        DNC C2002-131331
\mathsf{D}\mathsf{N}\mathsf{N}
     Administering and distributing substance, e.g. pharmaceutically active
{
m TI}
     agent, to target through bloodstream of organism by monitoring blood flow
     parameter(s), and adjusting distribution parameter.
     A96 B05 B07 P31 S03 S05
DC
     KENSEY, K
IN
     (KENS-I) KENSEY K; (RHEO-N) RHEOLOGICS INC
PA
CYC
     97
PI
     US 2002032149
                     A1 20020314 (200249) *
                                                 46
                     A2 20021010 (200277)
     WO 2002079778
                                           EN
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
            NL OA PT SD SE SL SZ TR TZ UG ZW
         W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK
            DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR
            KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PH PL PT RO
            RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG UZ VN YU ZA ZW
     AU 2002306461
                     A1 20021015 (200432)
ADT US 2002032149 A1 CIP of US 1997-919906 19970828, CIP of US 1999-439795
     19991112, CIP of US 2000-501856 20000210, CIP of US 2000-628401 20000801,
     CIP of US 2000-727950 20001201, CIP of US 2001-819924 20010328, US
     2001-841389 20010424; WO 2002079778 A2 WO 2002-US3984 20020207; AU
     2002306461 A1 AU 2002-306461 20020207
FDT US 2002032149 A1 CIP of US 6019735, CIP of US 6322524, CIP of US 6322525;
     AU 2002306461 A1 Based on WO 2002079778
PRAI US 2001-841389
                          20010424; US 1997-919906
                                                          19970828;
                          19991112; US 2000-501856
     US 1999-439795
                                                          20000210;
                          20000801; US 2000-727950
     US 2000-628401
                                                          20001201;
                          20010328; US 2001-828761
    US 2001-819924
                                                          20010409;
    US 2001-839785
                          20010420
    US2002032149 A UPAB: 20040520
AB
    NOVELTY - A substance (I) is administered and distributed (to a target)
     through a bloodstream of an organism by monitoring a blood flow
```

parameter(s) of the bloodstream, after which a distribution parameter is adjusted by altering the parameter(s).

DETAILED DESCRIPTION - A substance (I) is administered and distributed (to a target) through a bloodstream of an organism by monitoring a blood flow parameter(s) of the bloodstream, after which a distribution parameter is adjusted by altering the parameter(s). The parameter is circulating blood, absolute, effective, low shear or high shear viscosities, shear rate of circulating blood, work of heart, contractility of heart, thrombogenicity, platelet aggregation, lubricity, red blood cell deformability, thixotropy, yield stress, coagulability, coagulation time, agglutination, clot retraction, clot lysis time, sedimentation rate, or prothrombin rate.

USE - The method is used for distributing and administering a substance, e.g. pharmaceutically active agent, through a bloodstream of an organism such as a human. It is used for utilizing the viscosity of the circulating blood of a living being, for diagnostics and treatment.

ADVANTAGE - The method provides data in a short span of time, with minimal invasiveness, and without the need to directly measure pressure, flow, and volume.

Dwg.0/22

TECH

UPTX: 20020802

TECHNOLOGY FOCUS - INSTRUMENTATION AND TESTING - Preferred Components: The target is a cell, tissue or a system. The blood flow parameter is selected from intravenous diluents, red blood cell deformability agents, antiurea agents, oral contraceptives, anti-diabetic agents, antiarrythmics, antihypertensives, antihyperlipidemics, antiplatelet agents, anti-coagulants, appetite suppressants, antiobesity agents, blood modifiers, smoking deterrent agents, nutritional supplements, cholesterol-lowering agents, triglyceride-lowering agents, lubricants, homocysteine-reducing agents, vitamin supplements, beta-blockers, calcium channel blockers, ACE inhibitors, ACE-II inhibitors, vasodilators, blood pressure reducing agents, viscosity reducing agents, contractility reducing agents, hemodilution agents, adhesiveness minimizing agents, peripheral antiadrenergic/sympatholytics, anti-thrombogenic agent, warfarin, heparin, surfactants, saponifying agents, sodium bentonite magma, colloidal clays, colloidal silicon dioxide, micro-crystalline cellulose, gels of colloidal clays such as sodium bentonite, gels of organic polymers such as gelatin, agar, pectin, methylcellulose, or high-molecular-weight polyethylene glycol. TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Component: (I) (1 - 100 wt.%) is a pharmaceutically active agent selected from levonorgestrel, estrogen, progestin, (ethinyl) estradiol, ethynodiol, medroxyprogesterone, desogestrel, cyproterone, norethindrone, gestodene, norgestrel, mestranol, norgestimate, metformin, acarbose, insulin, chlorpropamide, glipizide, glyburide, tolazamide, glimepiride, troglitazone, pioglitazone, repaglinide, losartan potassium, candesartan cilexetil, irbesartan, mitiglinide, trendolapril/verapamil, nateglinide, nifedipine, nisoldipine, nicardipine, bepridil, isradipine, nimodipine, felodipine, amlodipine, diltiazem, verapamil, isosorbide, mononitrate, isosorbide dinitrate, nitroglycerin, hydralazine, minoxidil, hydrochlorothiazide, chlorothiazide, indapamide, metolazone, furosemide, bumetanide, ethacrynic acid, torsemide, spironolactone, triamterene, acetazolamide, mannitol, atenolol, bisoprolol, pindolol, metoprolol, timolol, nadolol, propanolol, carvedilol, captopril, fosinopril, benazepril, lisinopril, perindopril, enalapril, quinapril, losartan, valsartan, eprosartan, trandorapril, fenoldopam, ramipril, doxazosin, milrinone, benidipine, lemakalim, fantofarone, lemildipine, pirmenol, clentiazem, nebivolol, oxodipine, sematilide, pranidipine, nifekalant, aranidipine, barnidipine, lacidipine,

bucindolol, azelnidipine, dofetilide, ibutilide, watanidipine,

lercanidipine, landiolol, telmisartan, furnidipine, azimilide, CHF 1521, valsartan/hydrochlorothlazide, enalapril/nitrondipine, sotalol, arbutamine, olmesartan, conivaptan, sumatriptan, milrinone, lovastatin, atorvastatin, cerivastatin, simvastatin, fluvastatin, cholestyramine, colestipol, clofibrate, gemfibrosil, fenofibrate, pamaqueside, pitavastatin, phentermine, phendimetrazine, sibutramine, orlistat, aspirin, warfarin, enoxaparin, heparin, low molecular weight heparin, cilostazol, clopidogrel, ticlopidine, tirofiban, abciximab, dipyridamole, plasma protein fraction, human albumin, low molecular weight dextran, hetastarch, reteplase, alteplase, streptokinase, urokinase, dalteparin, filgrastin, immunoglogulin, ginkolide B, hirudins, foropafant, rocepafant, bivalirudin, dermatan sulfate mediolanum, eptilibatide, thrombomodulin, low molecular weight dermatan sulfate-opocrin, eptacog alfa, argatroban, fondaparinux sodium, tifacogin, lepirudin, desirudin, OP2000, melagatran, roxifiban, parnaparin sodium, human hemoglobin (Hemosol), bovine hemoglobin (Biopure), human hemoglobin (Northfield), antithrombin III, RSR 13, heparin-oral (Emisphere) transgenic antithrombin III, H37695, mesoglycan, CTC111, nicotine, buprorion, fasudil, ziconotide, amino acid preparations, minerals, electrolytes, vitamins, calcitriol, terbinafine, ticarcillin disodium, cefixime, meropenem, cefprozil, levofloxacin, cefpodoxime proxetil, imipenem, cefuroxime axetil, trovafloxacin, mupirocin, stavudine, didanosine, nevirapine, lamivudine, zidovudine, valcyclovir, ganciclovir, nefiracetam, remifentanil, sevoflurane, tiagabine, topiramate, lamotrigine, naratriptan, bromocriptine, tolcapone, oxaprozin, diclofenac, misoprostol, nabumetone, granisetron, dotarizine, RSR13, zonisamide, BMS204352, oxcarbazepine, tropisetron, irinotecan, topetecan, anastrozole, nilutamide, cladribine, gemcitabine, letrozole, vinorelbine, epirubicin, raloxifene, calcitonin, somatotropin, recombinant somatotropin, tolterodine, temiverine, meluadrine tartrate, lansoprazole, ropivacaine, bambuterol, israpafant, rupatadine, levosalbutamol, ARC68397AA, salbutamol (powder), salbutamol (inhalation), salbutamol (oral), salbutamol (powder inhalation), formoterol, salmeterol/fluticasone propionate, salmeterol MDI dose counter, salmeterol (inhalation), salmeterol hydrofluoroalkane, budesonide/formoterol, olopatadine, levobetaxolol, levobunolol, latanoprost/timolol, ketotifen, desferoxamine, leukine, sargramostin, or GM-CSF.

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ANSWER 13 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN 2002-372354 [40] WPIDS

DNC C2002-105446

TI Production of nateglinide B-form crystals containing no H-form crystals, by drying wet crystals of nateglinide solvate at low temperature until solvent disappears
```

DC B05
IN MARUO, M; MATSUZAWA, Y; MIYAZAKI, K; NISHINA, S; SUMIKAWA, M
PA (AJIN) AJINOMOTO CO INC; (AJIN) AJINOMOTO KK

CYC 98

PI WO 2002034713 A1 20020502 (200240)\* JA 9
RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PH PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001096001 A 20020506 (200257) EP 1334964 A1 20030813 (200355)

and performing crystal transformation.

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR

EN

KR 2003059212 A 20030707 (200377)

US 2003229249 A1 20031211 (200382)

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BR 2001014846 A 20040225 (200416)
     JP 2002537707 X 20040304 (200417)
     MX 2003003575 A1 20030701 (200423)
                     A 20040317 (200437)
     CN 1483018
ADT WO 2002034713 A1 WO 2001-JP9293 20011023; AU 2001096001 A AU 2001-96001
     20011023; EP 1334964 A1 EP 2001-976819 20011023, WO 2001-JP9293 20011023;
     KR 2003059212 A KR 2003-705671 20030424; US 2003229249 A1 Cont of WO
     2001-JP9293 20011023, US 2003-421888 20030424; BR 2001014846 A BR
     2001-14846 20011023, WO 2001-JP9293 20011023; JP 2002537707 X WO
     2001-JP9293 20011023, JP 2002-537707 20011023; MX 2003003575 A1 WO
     2001-JP9293 20011023, MX 2003-3575 20030423; CN 1483018 A CN 2001-821299
     20011023
FDT AU 2001096001 A Based on WO 2002034713; EP 1334964 A1 Based on WO
     2002034713; BR 2001014846 A Based on WO 2002034713; JP 2002537707 X Based
     on WO 2002034713; MX 2003003575 A1 Based on WO 2002034713
PRAI JP 2000-324375
                          20001024
     WO 200234713 A UPAB: 20020626
AΒ
     NOVELTY - Production of nateglinide (N-(trans-4-isopropyl-
     cyclohexane carbonyl)-D-phenylalanine) B-form crystals
     containing no H-form crystals, comprises drying wet
     crystals of nateglinide solvate at a low temperature
     until the solvent disappears and performing crystal
     transformation.
          ACTIVITY - Antidiabetic.
          MECHANISM OF ACTION - None given.
          USE - The nateglinide B-form crystals containing
     no H-form crystals are used as diabetes medicines.
          ADVANTAGE - The nateglinide B-form crystals
     containing no H-form crystals can be produced on an industrial
     scale at low cost.
     Dwg.0/0
TECH
                    UPTX: 20020626
     TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Method: No H-form
     crystals are detected by DSC. Crystallization is
     performed at at most50degreesC. The crystal transformation is
     performed by heating to 60-110degreesC. Both processes of the low
     temperature drying and the crystal transformation are processes
     which are performed on an industrial scale.
     ANSWER 14 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN
L3
AN
     2002-372336 [40]
                        WPIDS
DNC
     C2002-105445
     New composition comprises nateglinide in the amorphous state,
{
m TI}
     useful for treatibg diabetes.
DC
     B05
     MAKINO, C; NINOMIYA, N; YABUKI, A
IN
     (AJIN) AJINOMOTO CO INC; (AJIN) AJINOMOTO KK
PA
CYC
     98
PI
                    A1 20020502 (200240) * JA
     WO 2002034254
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
           NL OA PT SD SE SL SZ TR TZ UG ZW
        W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ DE DK
           DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR
           KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PH PL PT RO
           RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
                    A 20020506 (200257)
    AU 2001095999
    EP 1334720
                    A1 20030813 (200355)
                                          EN
        R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT
           RO SE SI TR
```

KR 2003042027 A 20030527 (200361)

AB

L3

AN

CR

 ${f T}{f I}$ 

DC

IN

PA

CYC

 $\mathtt{PI}$ 

DNC

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BR 2001014896 A 20030812 (200367)
     US 2004014815 A1 20040122 (200407)
     CN 1482903
                    A 20040317 (200437)
     JP 2002537308 X 20040826 (200456)
ADT WO 2002034254 A1 WO 2001-JP9291 20011023; AU 2001095999 A AU 2001-95999
     20011023; EP 1334720 A1 EP 2001-976817 20011023, WO 2001-JP9291 20011023;
     KR 2003042027 A KR 2003-705634 20030423; BR 2001014896 A BR 2001-14896
     20011023, WO 2001-JP9291 20011023; US 2004014815 A1 Cont of WO 2001-JP9291
     20011023, US 2003-421898 20030424; CN 1482903 A CN 2001-821217 20011023;
     JP 2002537308 X WO 2001-JP9291 20011023, JP 2002-537308 20011023
FDT AU 2001095999 A Based on WO 2002034254; EP 1334720 A1 Based on WO
     2002034254; BR 2001014896 A Based on WO 2002034254; JP 2002537308 X Based
     on WO 2002034254
PRAI JP 2000-324373
                          20001024
    WO 200234254 A UPAB: 20020626
    NOVELTY - Composition comprising nateglinide in the amorphous
     state, is new.
         ACTIVITY - Antidiabetic. In oral bioavailability studies in beagles
    amorphous nateglinide had an AUC ( mu g/ml.hr) of 22.29, a Cmax
     ( mu g/ml) of 9.46 and a Tmax (hr) of 0.38. The corresponding values for
    nateglinide crystalline form H were 20.53, 8.93 and 0.38
    respectively.
         MECHANISM OF ACTION - None given.
         USE - As preparations for administering nateglinide useful
    as an antidiabetic agent.
         ADVANTAGE - Have rapid release properties.
    Dwq.0/9
TECH
                   UPTX: 20020626
    TECHNOLOGY FOCUS - ORGANIC CHEMISTRY - Preferred Composition: The
    composition comprises amorphous nateglinide prepared by removing
    the solvent (preferably aqueous ethanol) from a solution of
    nateglinide and a hydrophiliic compound (preferably a water
    soluble polymer, water swellable polymer, sugar alcohol or salt,
    especially methylcellulose SM-4, hydroxypropylcellulose SL,
    hydroxypropylcellulose SSL, polyethylene glycol, sorbitol, xylitol,
    mannitol or crospovidone).
    ANSWER 15 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN
    2001-290407 [30]
                       WPIDS
    2003-401332 [38]
    C2001-088908
    Use of a combination of nateglinide with another antidiabetic
    compound for treating a metabolic disorder, e.g. diabetes and associated
    conditions, or for effecting weight loss.
    A96 B05
    ALLISON, M; GATLIN, M R; GUITARD, C; KARNACHI, A A; MANNION, R O;
    PONGOWSKI, M; BALL, M; KAMACHI, A A; BALL, M A
    (NOVS) NOVARTIS AG; (NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH; (ALLI-I)
    ALLISON M; (BALL-I) BALL M A; (GATL-I) GATLIN M R; (GUIT-I) GUITARD C;
    (KARN-I) KARNACHI A A; (MANN-I) MANNION R O
    95
                    A2 20010329 (200130) * EN
    WO 2001021159
                                               60
       RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ
           NL OA PT SD SE SL SZ TZ UG ZW
        W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM
           DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC
           LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE
           SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW
                    Al 20010323 (200130)
    FR 2798592
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FI 2001000683
                     A 20010515 (200140)
     AU 2000079044 A 20010424 (200141)
     CZ 2001001723 A3 20010815 (200157)
     MX 2001004255
                     A1 20010801 (200238)
                     A2 20020612 (200239)
     EP 1212077
                                           EN
         R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT
            RO SE SI
     NO 2002001197 A 20020516 (200240)
                     A 20020611 (200248)
     BR 2000014525
                     A3 20020702 (200253)
     SK 2002000360
                    A5 20020702 (200257)
     BE 1013726
                     A 20020523 (200274)
     KR 2002038758
     JP 2003509457 W 20030311 (200319)
                                                83
     US 2003162816 A1 20030828 (200357)
     NZ 517280
                    A 20040227 (200418)
     ZA 2002002107 A 20040331 (200426)
                                                86
     HU 2004000932 A2 20040728 (200454)
   WO 2001021159 A2 WO 2000-EP9074 20000915; FR 2798592 A1 FR 2000-11782
ADT
     20000915; FI 2001000683 A WO 2000-EP9074 20000915, FI 2001-683 20010402;
     AU 2000079044 A AU 2000-79044 20000915; CZ 2001001723 A3 WO 2000-EP9074
     20000915, CZ 2001-1723 20000915; MX 2001004255 A1 MX 2001-4255 20010427;
     EP 1212077 A2 EP 2000-969260 20000915, WO 2000-EP9074 20000915; NO
     2002001197 A WO 2000-EP9074 20000915, NO 2002-1197 20020311; BR 2000014525
     A BR 2000-14525 20000915, WO 2000-EP9074 20000915; SK 2002000360 A3 WO
     2000-EP9074 20000915, SK 2002-360 20000915; BE 1013726 A5 BE 2000-585
     20000915; KR 2002038758 A KR 2002-703551 20020316; JP 2003509457 W WO
     2000-EP9074 20000915, JP 2001-524585 20000915; US 2003162816 A1
     Provisional US 1999-240911P 19990917, Provisional US 2000-240918P
     20000309, Provisional US 2000-304196P 20000407, Cont of US 2000-663264
     20000915, US 2003-345908 20030116; NZ 517280 A NZ 2000-517280 20000915, WO
     2000-EP9074 20000915; ZA 2002002107 A ZA 2002-2107 20020314; HU 2004000932
    A2 WO 2000-EP9074 20000915, HU 2004-932 20000915
FDT AU 2000079044 A Based on WO 2001021159; CZ 2001001723 A3 Based on WO
     2001021159; EP 1212077 A2 Based on WO 2001021159; BR 2000014525 A Based on
     WO 2001021159; SK 2002000360 A3 Based on WO 2001021159; JP 2003509457 W
    Based on WO 2001021159; NZ 517280 A Div in NZ 528738, Based on WO
     2001021159; HU 2004000932 A2 Based on WO 2001021159
                         20000826; US 1999-398364
PRAI GB 2000-21055
                                                         19990917;
     US 2000-545480
                         20000407
AB
     WO 200121159 A UPAB: 20040823
    NOVELTY - Nateglinide (I), optionally in combination with
    another antidiabetic compound, can be used in the treatment of diabetes
    and associated conditions. The combination can also be used for effecting
    weight loss.
         DETAILED DESCRIPTION - Use of a combination of nateglinide
     (I) and at least 1 other antidiabetic compound, selected from thiazolidine
    derivatives (glitazones), sulfonyl urea derivatives and metformin, present
    in the free form or as salts, for prevention, delay of progression or
    treatment of metabolic disorders, or for cosmetic treatment to effect a
    loss of body weight, is new.
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INDEPENDENT CLAIMS are included for the following:

- (a) a combination of (I) with an antidiabetic compound (as described above) for simultaneous, sequential or separate use;
  - (b) compositions comprising (I) with the antidiabetic compound; and
- (c) a composition capable of being granulated in the presence of water without the need for a subsequent pulverization step prior to tabletting, comprising (I) and a carrier; and its use for treating a metabolic disorder.

ACTIVITY - Antidiabetic; anorectic; antilipemic; opthalmological; vasotropic; antiulcer; antiinflammatory; cardiant; hypotensive;

antianginal; dermatological; antiarthritic; osteopathic; gastrointestinal. MECHANISM OF ACTION - None given.

USE - For treating a metabolic disorder, e.g. diabetes (particularly type II diabetes mellitus) and associated conditions, also for effecting weight loss. The compositions can be used to treat e.g. hyperglycemia, hyperinsulinemia, hyperlipidemia, insulin resistance, impaired glucose metabolism, obesity, diabetic retinopathy, macular degeneration, cataracts, diabetic nephropathy, glomerulonephritis, diabetic neuropathy, erectile dysfunction, premenstrual syndrome, vascular restenosis, ulcerative colitis, coronary heart disease, hypertension, angina pectoris, myocardial infarction, stroke, skin and connective tissue disorders, foot ulcerations, metabolic acidosis, arthritis, osteoporosis, and conditions of impaired glucose tolerance.

Dwg.0/0

TECH

UPTX: 20010603

TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Compounds: (I) is present in the B-type or H-type **crystal** modification. The antidiabetic compound is preferably a glitazone, e.g. rosiglitazone, troglitazone or pioglitazone, or metformin or its hydrochloride. Preferred Combination: The combination may further comprise insulin, or comprises at least 2 antidiabetic compounds.

Preferred Composition: A composition comprising (I) and a carrier releases 60-95 wt.% (I) within 30 minutes in water. The composition may further comprise colloidal silicon dioxide, and a disintegrant, preferably having molecular weight greater than 1000000, and particle size distribution of less than 400 microm or less than 74 microm. The composition may be in the form of a tablet, a granular composition, or contained in a capsule.

- L3 ANSWER 16 OF 16 WPIDS COPYRIGHT 2004 THOMSON DERWENT on STN
- AN 2001-281809 [29] WPIDS

DNC C2001-085742

- TI Combination used for treating diabetes and metabolic disorders comprises nateglinide, antidiabetic phenylacetic acid derivative or acarbose and carrier.
- DC B05
- IN BALL, M; DUNNING, B; GATLIN, M R; PONGOWSKI, M
- PA (NOVS) NOVARTIS AG; (NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH

CYC 95

- PI WO 2001026639 A2 20010419 (200129) \* EN 28
  - RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW
    - W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001011339 A 20010423 (200147)

EP 1218015 A2 20020703 (200251) EN

- R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI
- ADT WO 2001026639 A2 WO 2000-EP9816 20001006; AU 2001011339 A AU 2001-11339 20001006; EP 1218015 A2 EP 2000-972695 20001006, WO 2000-EP9816 20001006
- FDT AU 2001011339 A Based on WO 2001026639; EP 1218015 A2 Based on WO 2001026639

PRAI US 1999-415308 19991008; US 1999-415307 19991008

AB WO 200126639 A UPAB: 20010528

NOVELTY - Combination (I) comprises nateglinide, an antidiabetic phenylacetic acid derivative or acarbose or their salts and optionally at least one carrier for simultaneous, separate or sequential use.

DETAILED DESCRIPTION - An INDEPENDENT CLAIM is included for a commercial package comprising (I) together with instructions for the delay

of progression or treatment of metabolic disorders or a method of improving bodily appearance.

ACTIVITY - Antidiabetic; antilipemic; antilulcer; antiinflammatory; vasotropic; hypotensive; cardiant; antiarthritic; osteopathic; cerebroprotective; anorectic; gastrointestinal; ophthalmological; muscular; dermatological.

MECHANISM OF ACTION - None given.

USE - Used for treating diabetes, conditions associated with diabetes, especially type 2 diabetes mellitus and metabolic disorders e.g. hyperglycemia, hyperinsulinaemia, hyperlipidemia, insulin resistance, impaired glucose metabolism, obesity, diabetic retinopathy, macular degeneration, cataracts, diabetic nephropathy, glomerulosclerosis, diabetic neuropathy, erectile dysfunction, premenstrual syndrome, vascular restenosis and ulcerative colitis, coronary heart disease, hypertension, angina pectoris, myocardial infarction, stroke, skin, connective tissue disorders, foot ulcerations, metabolic acidosis, arthritis, osteoporosis and conditions of impaired glucose tolerance.

ADVANTAGE - The **nateglinide** and phenylacetic acid derivative show a synergistic effect.

Dwg.0/0

TECH

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UPTX: 20010528

TECHNOLOGY FOCUS - PHARMACEUTICALS - Preferred Combination: The combination is a combined preparation or a pharmaceutical composition. The antidiabetic phenylacetic acid is repaglinide or its salts. The combination also comprises at least one antidiabetic thiazolidinedione, sulfonyl urea derivatives, metformin or insulin or their salts or at least one further antidiabetic phenylacetic acid derivative or its salts. The nateglinide is present in the B-type or H-type crystal modification.